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THE ROLE OF BULK ADDITIONS IN SOLID LUBRICANT COMPACTS

PHILLIP W. CENTERS Lubrication Branch Fuels and Lubrication Division

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19. ABSTRACT (Continued)

 $Sb_2\theta_3$, or formation of a eutectic of antimony and molybdenum oxides to improve transfer film formation are refuted.

Evaluation of oxides and sulfides a bulk additions in MoS₂ indicate that the role of antimony trioxide is not unique; e.g., MoO_3 is superior to Sb_2O_3 at test temperature of 316°C. Confirmation is also provided by the finding that lower test temperatures are required for antimony oxide additions to graphite to improve performance of graphite, evidently resulting from higher asperity contact temperatures generated in the graphite compact.

A strong correlation of solid lubricant compact wear volume and weak correlation of friction coefficient with shear strength of additives are found. Therefore, the improvement in tribological performance is a bulk, rather than interfacial phenomenon.

In summary, lubricant compact properties and selection of bulk additions for improved tribological performance are hypothesized to be determined by the test or application conditions, the shear strength at asperity contact temperature of the additive and properties of the solid lubricant.

PREFACE

This technical report was prepared for the Lubrication Branch, Fuels and Lubrication Division, Aero Propulsion Laboratory (APL), Air Force Wright Aeronautical Laboratories (AFWAL), Air Force Systems Command, Wright-Patterson Air Force Base, Ohio. The work herein was accomplished under Project 3048, Task 304806, Work Unit 30480626, "Turbine Engine Lubricant Research" for the period November 1983 to November 1986, with Mr Phillip W. Centers, AFWAL/POSL, as project engineer. Special acknowledgement is given to Captain F. Dean Price, Chris Klenke, Robert Wright, Gayle Byrd and Corliss Goldsmith without whose assistance, this project could not have been completed.

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The author also wishes to express very special thanks to his mother, brother Bob, and his wife Connie for their love, patience and understanding.

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CHAPTER I

INTRODUCTION

An effort to develop high temperature solid lubricants for 315-540°C usage has recently been reported. (1) One potential application for such materials is to lubricate roller bearings in small turbine engines. (2) Typically, such an engine weighs about 65 kg, produces 3.3 kN thrust and has a shaft speed of up to 63,500 rpm. By use of solid rather than liquid lubrication, it may be possible to reduce engine weight and complexity while permitting an increase in thermodynamic efficiency due to higher operating temperature limits.

of lubricant candidates evaluated for the proposed application, a compact of molybdenum disulfide and antimony trioxide first described by Murphy, (3) and pursued by others, (4-6) appears to be most promising. Several investigators found that such compacts, whether evaluated with or without resinous binder, are superior to molybdenum disulfide when evaluated at temperatures up to 232°C, i.e., Lavik, Hubbell and McConnell (7) noted that pellets of MoS₂-Sb₂O₃ as well as bonded film formulations containing MoS₂-Sb₂O₃ evaluated in a standard pin in V-block configuration and in a dual rub-shoe test, were all

superior to MoS₂ without the additions. Those authors offered an explanation of the beneficial interaction by attributing the superior performance of MoS₂-Sb₂O₃ compacts to a hypothesized eutectic formed of MoO₃ and Sb₂O₃. It was proposed that the low melting eutectic reduces both friction and wear volume by enhancement of effective transfer film formation in both vacuum and air and reduces the detrimental effects of wear debris generated on sliding MoS₂ in air.

Subsequently, Harmer and Pantano⁽⁸⁾ studied MoS₂-Sb₂O₃ compacts. In that effort, chemical and mechanical properties were examined. They concluded that (i) Sb₂O₃ influenced the oxidation of MoS₂, (ii) Sb₂O₃ or Sb₂O₄ reacted with MoO₃ to produce a dense product, (iii) the morphology of wear surfaces are different if compacts contain Sb₂O₃ as compared with non-additive MoS₂ pellets, and (iv) Sb₂O₃ evidently modified the chemistry of burnished surfaces. The authors proposed that several areas were worthy of additional investigation but did not provide a conclusive explanation for the beneficial effects of Sb₂O₃ additions to MoS₂.

Nosov⁽⁹⁾ then evaluated MoS_2 , $MoS_2-Sb_2O_3$, and $MoS_2-Sb_2S_3$ solid lubricants in a dual roller test arrangement. Additions of Sb_2O_3 were found to increase MoS_2 lubricant life under both rolling and sliding conditions. If a roller coated with $MoS_2-Sb_2O_3$ were

pretreated by holding it at 350°C for three hours and then evaluated under ambient conditions, its life was typically increased significantly compared with MoS₂ identically pretreated. However, non-pretreated MoS₂-Sb₂O₃ formulations routinely only had 0-50% of the life of non-pretreated formulations. Nosov concluded that Sb₂O₃ and Sb₂S₃ improved the lubricating properties of MoS₂. Nosov speculated that the formation of a low melting eutectic proposed earlier ⁽⁷⁾ was not probable under the test conditions; however, it was stated that Sb₂O₃ increased the exidation resistance of MoS₂. Nosov did not offer supporting experimental evidence for either speculation.

Later, Gardos and McConnell⁽¹⁾ in their study on high-temperature, high-load lubricating composites evaluated at 316°C (600°F) found that the MoS₂-Sb₂O₃ formulation was comparable with, if not superior to, other candidate solid lubricants. Those authors speculated that Sb₂O₃ might act sacrificially in oxidizing to Sb₂O₄ or higher oxides, thus retarding MoS₂ oxidation, and providing improved high temperature compact properties.

In a recent study (10) of anti-galling agents for threaded parts, it was found that formulations containing MoS_2 , Sb_2O_3 , and a resinous binder were superior in rider-plate and bolt-nut tests. Those authors assumed that one role of Sb_2O_3 was to prevent MoS_2 oxidation. However,

those workers did not provide experimental data to support that explanation.

Thus, in summary, since a patent in $1967^{(3)}$ concerning the use of $\mathrm{Sb}_2\mathrm{O}_3$ in MoS_2 solid lubricant formulations, studies of friction and wear of $\mathrm{MoS}_2\mathrm{-Sb}_2\mathrm{O}_3$ compacts containing up to 45 weigh; percent $\mathrm{Sb}_2\mathrm{O}_3$ have demonstrated that such solid lubricants are superior to MoS_2 under a variety of test conditions. During that same period, no experimental study has been conducted to confirm or disprove any of a number of hypotheses advanced to explain the beneficial effect of $\mathrm{Sb}_2\mathrm{O}_3$ additions to MoS_2 .

Although the emphasis of most efforts hypothesize that the beneficial interaction of $\mathrm{Sb}_2\mathrm{O}_3$ and hoS_2 involves reduction of MoS_2 oxidation, many authors $^{(7-10)}$ describe the lubricating films formed when $\mathrm{Sb}_2\mathrm{O}_3$ is present as being more dense and sintered in appearance. None of the authors attribute the improved performance to the generation of a superior tribological surface by addition of $\mathrm{Sb}_2\mathrm{O}_3$.

Thus, several studies have concluded that formulations of $MoS_2-Sb_2O_3$ are superior to MoS_2 as solid lubricants. Alternative hypotheses have been made to rationalize the beneficial effects of Sb_2O_3 additions to MoS_2 ; however, a comprehensive study has not been undertaken to critically examine the validity of the hypotheses advanced and to elucidate the beneficial effects of bulk additions of Sb_2O_3

to MoS_2 or to determine if the beneficial mechanism is unique for solid lubricant compacts.

CHAPTER II

SCOPE OF THIS STUDY

Based upon a review of previous work regarding improved tribological properties of $MoS_2-Sb_2O_3$ formulations compared with MoS_2 solid lubricants, this effort was undertaken to critically examine the beneficial role played by Sb_2O_3 in MoS_2 solid lubricant compacts. That effort was organized as follows:

- (i) Study the kinetics of oxidation of So_2o_3 to determine if So_2o_3 plays an antioxidant xole in MoS_2 lubricant compacts.
- (ii) Examine the possibility of $Sb_2O_3-MoO_3$ or $Sb_2O_4-MoO_3$ eutectic formation and possible beneficial role for those eutectics.
- (iii) Mapping of the tribological characteristics of ${\rm Sb_2O_3-MoS_2}$ under various conditions to elucidate the performance of the system.
- (iv) Develop an understanding of the beneficial role of ${\rm Sb}_2{\rm O}_3$ and other additives in ${\rm MoS}_2$ and determine if that role is unique for ${\rm Sb}_2{\rm O}_3$ as an additive and for ${\rm MoS}_2$ as a solid lubricant.

(v) Develop a hypothesis for the behavior of bulk additions in solid lubricants to provide a basis for improvement of such systems.

CHAPTER III

LITERATURE SEARCH

Important in the study of beneficial bulk additions to solid lubricants is an understanding of friction and wear, lubrication, and protective mechanisms for solid compact systems containing a proven beneficial additive, ${\rm Sb}_2{\rm O}_3$. PRICTION

The laws of friction as summarized by Clauss (11) are:

(i) Frictional resistance is proportional to the normal load on sliding surfaces:

 $F = \mu N$

where F = frictional resistance

 μ = coefficient of friction

N = normal force

 μ may be additionally specified as $\mu_{g},$ the static coefficient of friction experienced at sliding initiation, and $\mu_{k},$ the kinetic coefficient obtained during constant sliding velocity.

- (ii) Frictional resistance is proportional to the true area of contact, but not to the apparent area of contact between two bodies.
- (iii) Frictional resistance is not proportional to velocity. There is an effect; but over a wide range of

sliding velocities, the coefficient tends to be rather stable unless frictional heating is involved.

(iv) The materials in contact do strongly affect frictional resistance, i.e., frictional coefficients may range from less than 0.02 to more than 1.0 depending upon the materials selected.

Additional considerations are that all surfaces are rough when examined in close detail. Early theorists thought that frictional work results from raising asperities over opposite asperities, and friction itself involves interlocking of opposing asperities. Thus, lubricants performed the simple task of filling voids between asperities and finally providing a smooth surface. However, modern theory claims that friction originates from two causes: shearing of welded junctions formed at asperity contact and ploughing that occurs as a harder material comes into contact with a softer one.

Clause also states that important considerations regarding friction are:

with the projected area of contact is very small compared with the projected area of contact. The amount of true contact may be roughly estimated by dividing the load by the yield strength of the material. That is, if the yield strength is 3.45 GPa and the load is 0.345 GPa, then a rough estimate of the fraction of real contact area compared to projected area is 0.10.

- (ii) High temperatures, e.g., 1250 K, (12) in the contact area may be generated under sliding even at ambient temperature.
- (iii) Finally, sliding is a "stick-slip" process.

 Under reseasable conditions, the process proceeds without detection by the casual observer.

WEAR

Wear, defined as the removal of material from solid surface, has been conveniently categorized by Rabinowicz⁽¹³⁾ as adhesive, abrasive, corrosive, surface fracture, erosive and fretting.

Adhesive wear is thought to be most common and originates in the attractive atomic forces as two surfaces are brought together. The adhesion may result in material removal from either surface dependent upon many material and environmental factors.

Quantitatively, it has been found that for adhesive wear:

- (i) Wear is directly proportional to load
- (ii) Wear is directly proportional to sliding distance
- (iii) Wear is typically inversely proportional to the

An achesive law of wear as developed by Archard (14) is as follows:

$$V = \frac{k L x}{3p} \tag{3.1}$$

where V = wear volume

L = load

x = sliding distance

p = hardness of material being worn

k = material wear constant

Thus, it is possible to predict the effect of changing critical parameters in a typical wear experiment. The relationship does not consider the effects in changes in temperature or environment that occur in wear situations.

The other common form of wear is abrasive. This form of wear occurs when a hard surface slides across a softer one creating grooves with the loss of material. Abrasive wear can be differentiated as two- or three-body wear. Two-body wear occurs when a hard, rough surface interacts with another surface, while three-body wear occurs when hard, abrasive particles interact at the sliding interface of two other materials.

A simple abrasive wear equation proposed by Rabinowicz (13) which takes the same form as the equation for adhesive wear is as follows:

$$V = \frac{\tan \theta \ Lx}{\pi \ p} \tag{3.2}$$

where V, L and x are as defined in the adhesive equation and

p = hardness of softer surface

 $\tan \theta$ = weighted average of $\tan \theta$'s for all abrasive cones where θ is the angle between the intersecting cone and the surface minus 180°. Rabinowicz' equation yields an abrasive wear constant, k_{abr} , as follows:

$$k_{abr} = 0.96 \overline{\tan \theta} \tag{3.3}$$

Thus, it is possible to generate similar, simplified relationships for adhesive and abrasive wear.

For other less common types of wear, which should not be significant in this effort, similar mathematical relationships have been derived which may be applied for engineering purposes. (13)

LUBRICATION

The most effective method of lubrication (15) is to provide a hydrodynamic film between two surfaces, in which a fluid film is drawn into the contact area separating two sliding surfaces. The coefficient of friction is very low, and with good design and maintenance, component lives can be exceedingly long. When the load becomes great, the lubrication regime is elastohydrodynamic. In that regime,

lubrication is influenced by the elastic properties of the substrates. Under very high load, surface asperities come into contact, and boundary lubrication begins. Boundary lubrication occurs frequently in high load sliding, e.g., autrmotive and truck differentials. That type of application requires an extreme pressure additive which provides a source of renewable surface boundary lubricant. The extreme pressure additive reacts at hot spots on the surface and forms a solid lubricating surface. For most lubrication applications, fluids are adequate and perform remarkably well especially when formulated with an additive package, e.g., anti-oxidants, dispersants, and/or viscosity improvers to enable their use over a wider temperature range. However, applications under extreme environmental conditions, such as very high temperatures or vacuum, preclude the use of fluid lubricants and extreme pressure additives. In these cases a solid lubricent may be singularly applied as a replacement for a liquid lubricant or combined with a solid organic resin to form a compact for high temperature applications.

SOLID LUBRICANTS

A solid lubricant has been defined as "any solid used between two surfaces to provide protection from damage during relative movements to reduce friction and wear". (16) Many references are available (16) in which such

materials are discussed at length. Some advantages cited for use of solid lubricants include good stability at extreme temperatures and in chamically active environments which permit greater flexibility in design. Solid lubricants can be used under extremely high loads, and they are lightweight compared with conventional liquid systems. Solid lubricants, however, have many disadvantages, including a higher coefficient of friction than that in hydrodynamic lubrication, unavoidable solid sliding contact wear, finite lifetime, and no cooling capability.

Solid lubricants may be classified into several categories: (i) soft metals, such as silver and gold; (ii) laminar solids, such as graphite, molybdenum disulfide, and tungsten disulfide; (iii) non-laminar solids, such as lead monoxide, silver iodide, and calcium fluoride; (iv) complex solid compacts, exemplified by tungsten diselenide/gallium/indium; (v) soaps, fats, and waxes, such as stearic acid; and (vi) polymers, such as polytetrafluoroethylene. Each of the above solid lubricants functions by a common mechanism within that category, but typically unlike those of different categories. Also, often it is the application that critically determines which class and even which specific solid lubricant may be used.

Properties of candidate solid lubricants related to friction and wear are shear strength, hardness, mating surface adhesion and ductility. Important propeties related to overall solid lubricant performance are thermal stability, chemical inertness, particle size, purity, surface mobility, corrosion prevention and electrical conductivity. These property relationships and interactions are often complex and lead to development of completely different capabilities even within categories. For example, graphite and molybdenum disulfide lubricate by shearing interlaminarly with small forces while carrying high normal loads. For graphite, it has been observed that adsorbed water and gases improve lubrication performance. However, adsorbed species are not required for molybdenum disulfide lubrication. (17) Therefore, exidatively stable graphite loses adsorbed species and, thus, much of its lubricating ability at higher temperatures, whereas MoS2 lubricates effectively to relatively high temperatures without adsorbed species, but oxidizes at lower temperatures than graphite. The comparison of graphite with MoS_2 demonstrates that within each category of solid lubricants there may be examples of dissimilar performance requiring knowledgeable assessment to match a solid lubricant with an application.

Not only do solid lubricants differ chemically, physically, and in mechanisms of lubrication, but also

application methods include dispersions in oils and greases, fine powders, organic monolayers and films established by transfer, burnishing, plating, sputtering, or resin-bonding. Although transfer and burnished films have been employed with limited success over a wide temperature range, resin-bonded systems are under intensive evaluation for high temperature appliations. (16)

High temperature properties of several solid lubricants have encouraged investigations of solid lubricant capabilities in serospace applications. Of current interest is the possibility of using solid lubricants in selected turbine engine mainshaft bearings for limited lifetime applications. (1) The rationale for using solid lubricants in such engines is that weight savings will result from the elimination of the liquid lubrication system. Also, higher operating temperatures permissible with the use of a solid lubricant shall permit improved thermodynamic efficiency. Additionally, the short operational time requirement minimizes problems resulting from the limited life of solid lubricants.

The potential for solid lubrication of small turbine engines has generated considerable developmental effort in the area of high temperature solid lubricants. (4) Many of recent developmental efforts have focused on molybdenum disulfide or other dichalcogenides. The potential

lubricating dichalcogenides are formed from Group IVB, VB, and VIB metals with a chalcogen, i.e., sulfur, selenium and tellurium. There are approximately sixty dichalcogenides formed, with two-thirds of these being layered. Of the possibilities, Magie, (18) Jamison, (19) and Gardos (1) have found that structural considerations limit good solid lubricating properties to the molybdenum, tungsten, and some niobium dichalcogenides. MoS₂ is the well-known example of this class.

Jamison⁽²⁰⁾ has observed that from a Madhukar plot⁽²¹⁾ of the ratio of cation/anion radii versus ionicity of the bond that good solid lubricants are to be found in one area of that plot. His data imply that there are basic structural and bonding parameters which are required for a dichalcogenide to be a useful solid lubricant. The data of Jamison and Cosgrove⁽²²⁾ was employed to correlate predicted and actual performance of the solid lubricants.

Also, it has been observed that good solid lubricants have a trigonal prism structure with a minimum c/a (unit cell dimension ratio) of 1.93, while poor lubricants have the trigonal antiprism structure with a c/a smaller than that of the good lubricants (e.g., 1.6 or near the hardsphere full density model). In solid lubrication studies, "c" is standardized by dividing the true crystallographic "c" value by the number of X-M-X layers in a single unit cell, where "X" is the chalcogen and "h" is the metal.

been made to prepare intercalates, i.e., materials with atoms inserted into the X-X layer. The intercalates have been predicted to have superior lubricating properties as compared with nonintercalates. The premise is that if atoms can be introduced between the chalcogenic layers, then shearing will require less force; thus, lubrication would be improved. (19)

Compacts consisting of $WS_2/Ga/In$ have also been developed (23,24) and have excellent static thermal properties and good room temperature lubricant properties. When compared with MoS_2 , however, the alternative solid lubricant may have one modestly superior property, but other properties, such as fatigue strength often fall short of that of MoS_2 . (25) These materials also must be produced synthetically at considerable cost while the natural MoS_2 ore, molybdenite, can be processed for use as a solid lubricant with little effort at lower costs. Thus, the search for a good high temperature solid lubricant for many applications has not proven truitful.

PROPERTIES OF AND OXIDATION OF MOLYBDENUM DISULFIDE

Historically, molybdenum disulfide $^{(26)}$ was confused with graphite because general appearance and lubricating characteristics were similar. That confusion ended when Scheele demonstrated in 1778 that MoS_2 contained sulfur and prepared molybdic acid $(MoO_3 \cdot 2H_2O)$ from molybdenite. $^{(27)}$

Molybdenum disulfide is a soft lead-gray compound. It has a metallic luster and contains 59.94 weight percent molybdenum and 40.06 weight percent sulfur. MoS_2 has a specific gravity of 4.8 and melts at $1185^{\circ}C.^{(28)}$ Other sulfides such as Mo_3S_4 , Mo_2S_3 , MoS_4 and MoS_3 are known. MoS_3 are known but are not stable. MoS_4

A comprehensive review⁽¹⁷⁾ of molybdenum disulfide structures and lubricating properties has been prepared. The structure was first determined by Dickinson and Pauling.⁽³⁰⁾ Molybdenum is strongly bonded between layers of sulfur while adjacent sulfur layers are only weakly bonded by van der Waals forces. Sliding takes place interlaminarly between sulfur layers and, unlike graphite, no interlaminar adsorption of water or condensable gases is required for lubricating action; however, water vapor does have a beneficial effect up to about 65% relative humidity.

The review⁽¹⁷⁾ also summarizes the effect of load, sliding speed, temperature, vacuum, radiation, substrate, particle size, impurities and crystallographic orientation on the behavior of MoS₂ as a lubricant. The substrate and orientation of MoS₂ films does affect frictional properties. Particle size effects are related to ease of oxidation and the formation of MoO₃.

Of the MoS_2 oxidation studies, Cardoen's (31) appears to be the most comprehensive. In that study, compacts of molybdenite were oxidized under various oxygen

concentrations. After test, weights of pellets and observation of the oxidation interface, if available, were used to assess oxidation behavior over the 450° to 700°C temperature range. In the studies, the formation of MoO_2 was considered and calculations were made which showed that MoO_2 is metastable at temperatures over 350°C and at oxygen pressures exceeding 10^{-21} atmosphere.

Cardoen⁽³¹⁾ was also able to define five oxidation regions for molybdenite. It should be emphasized that those regions were established for the oxidation of molybdenite containing impurities, not pure MoS₂, and for compressed samples in which cracks and pathways for oxygen transport probably exist. Both objections were addressed by Cardoen, although his arguments are not completely satisfying.

In discussion, Cardoen notes that there are five steps in a gas-solid reaction. Those steps are:

- (i) Diffusion of reactants to solid surface.
- (ii) / Adsorption of reactants on the surface.
- (iii) Reaction on the surface.
- (iv) Desorption of products of reaction.
- (v) Diffusion of products away from surface.
 If steps (i) or (v) are rate controlling, the rates are parabolic and usually involve a protective coating of reaction products.

The Pilling-Bedworth (PB) ratio⁽³²⁾ is used to assess the possibility of a protective layer being formed.

Essentially, if the ratio of the volume of oxide formed is greater than the volume of the solid from which it is formed, then the oxide may protect (PB = 1 to 2). However, if the PB is less than one, then coverage of the substrate cannot be complete and the outer layer will not protect.

Of course, if PB is large, then the layer can spall and provide reduced protection. The ratios are only

The PB for MoO₃/MoS₂ is 0.916. Therefore, Cardoen⁽³¹⁾ concluded that the oxide is nonprotective. This conclusion was supported by his experimental data since linear kinetics were observed. His linear rate law conclusion is unlikely the parabolic laws found by others.⁽³³⁾ Cardoen⁽³¹⁾ cite—the exothermic surface reaction as the origin of incorrect rate law interpretation problems of others.

Ba ad upon his results, Cardoen proposed that the linear exidation of molybdenite takes place as follows:

$$|s + o_2| \Rightarrow | + |s + o_2|$$

 $|s + o_2| + |s + o_2|$
 $|s + o_2|$

where represents the surface and + an activated complex.

Modern surface analytical instrumentation has been employed to examine molybdenum sulfide and molybdenum oxide surfaces. (34-42)

ANTIMONY OXIDES

Antimony $^{(43)}$ forms several oxides which are listed along with handbook data $^{(28)}$ on those oxides in Table 1. In the listed oxides, antimony is either tri or penta valent. $\mathrm{Sb}_2\mathrm{O}_4$ ($\mathrm{Sb}_2\mathrm{O}_3$ · $\mathrm{Sb}_2\mathrm{O}_5$) contains both tri and penta valent antimony. Additionally, evidence exists for $\mathrm{Sb}_6\mathrm{O}_{13}$. $^{(44)}$ hydrated forms of $\mathrm{Sb}_2\mathrm{O}_3$ and $\mathrm{Sb}_2\mathrm{O}_5$ are known as antimonious and antimonic acids, respectively. Antimony and some antimony compounds are toxic and must be handled cautiously.

Remy⁽⁴⁵⁾ reported that the cubic structure of the dimorphic trioxide is composed of Sb_40_6 groups (antimony sequioxide) occupying the points of a diamond lattice with a cube edge of 11.14Å. Adjacent molecular centers are 4.84Å apart. The Sb_40_6 molecule consists of an octahedron of oxygen atoms, with Sb atoms forming an interpenetrating tetrahedron. The Sb-0 distance is 2.22Å. The orthorhombic dimorph has infinite chains of Sb_20_3 composition where weak Sb-0 bonds hold the structure together.

Roberts and Fenwick⁽⁴⁶⁾ found that the trioxide in the low temperature cubic form is stable up to the transition temperature of 570 \pm 10°C, while the orthorhombic dimorph

TABLE 1. PROPERTIES OF ANTIHONY AND ANTIHONY OXIDES

ELEMENT OR COMPOUND		ORMULA EIGHT	CRYSTAL FORM	SP GRAV	MELT POINT (°C)	BOILING POINT (°C)
Antimony	Sb	121.75	Hex, Silver White Metal	6.69	630.5	1380
Senarmontite	sb ₂ 0 ₃ (c)	291.5	Cubic, White	5.2	656	1550 (subl)
Valentinite	sb ₂ 0 ₃ (o)	291.5	Rhomb. Colorle	5.67	656	1550
Cervantite	Sb ₂ 0 ₄	307.5	White Powder	5.82	-0,930	
Antimony Pentaoxide	sb ₂ 0 ₅	323.5	Yellow Powder	3.80	-0,380 -0,93	0

is stable up to the melting point of about $650\,^{\circ}\text{C}$. $\text{Sh}_2\text{O}_3(\circ)$ exists at room temperature in a metastable state.

More recent data generated by White, et al., $^{(47)}$ in a high pressure study of the transition temperature of $8b_2c_3$ indicates a reversible transition temperature of 606°C with a heat of transition of about 1000 cal/mole. They employed the Clausius-Clapeyron relationship as follows:

$$\frac{dp}{dT} = \frac{\Delta H}{T\Delta V} \tag{3.3}$$

where p = pressure

T = temperature

 ΔH = enthalpy of transition

 $\Delta V = volume change$

and calculated molar volumes from x-ray data of 52.216 cm³ for $Sb_2O_3(c)$ and 50.021 cm³ for $Sb_2O_3(c)$ to determine the heat of transition.

The metastable $\mathrm{Sb}_2\mathrm{O}_3(c)$ to $\mathrm{Sb}_2\mathrm{O}_3(c)$ transition has been studied in the $490\text{--}530\,^\circ\mathrm{C}$ temperature range. (48) In that effort, infrared spectroscopy was employed to analyze mixtures after thermal treatment. The activation energy for the transformation was found to range from 46 to 50.8 kcal/mole. It was claimed that heterogeneous nucleation of the $\mathrm{Sb}_2\mathrm{O}_3(c)$ on the $\mathrm{Sb}_2\mathrm{O}_3(o)$ surface is followed by growth of the more stable $\mathrm{Sb}_2\mathrm{O}_3(o)$. The rate controlling step was stated to be sublimation of $\mathrm{Sb}_2\mathrm{O}_3(o)$. Additional

supporting evidence was offered in noting that the heat of sublimation of $\mathrm{Sb}_2\mathrm{O}_3(o)$ is about 43.6 kcal/mole. This concept supported by the sublimation pressure data of Hincke⁽⁴⁹⁾ and Jungermann and Plieth.⁽⁵⁰⁾ Hincke⁽⁴⁹⁾ measured the vapor sublimation pressure data of the solid $\mathrm{Sb}_2\mathrm{O}_3$ dimorphs over the temperature range of 470° to 650°C by weighing sublimate in an inert atmosphere. The vapor pressure of liquid $\mathrm{Sb}_2\mathrm{O}_3$ was determined from 656° to 800°C. The vapor pressure as function of temperature relationships are expressed as follows:

$$\log_{10}p = 12.195 - \frac{10357}{T}$$
 (cubic) (3.4)

$$\log_{10}p = 11.318 - \frac{9625}{T}$$
 (orthorhombic) (3.5)

$$\log_{10}p = 5.137 - \frac{3900}{T}$$
 (liquid) (3.6)

where p = sublimation pressure (torr)
and T = absolute temperature (K).

The heats of sublimation found by Hincke are 47.32 kcal/mole (cubic), 44.08 kcal/mole (orthorhombic), and heat of vaporization is 17.83 kcal/mole (liquid). The heat of transition was found to be 3.24 kcal/mole at 557°C. Heat of fusion for the cubic form was found to be 29.49 kcal/mole and for the orthorhombic form, 26.25 kcal/mole.

Much more recently, Jungermann and Plieth $^{(50)}$ used a Knudsen cell and a sublimation weight apparatus to derive the following sublimation pressure equations:

$$\log_{10}p = 11.74 - \frac{9568}{T}$$
 (cubic) (3.7)

$$log_{10}p = 11.35 - \frac{9535}{T}$$
 (orthorhombic) (3.8)

where p and T are defined above.

Jungermann and Plieth found the heat of sublimation for $Sb_2O_3(c)$ is 43.8 \pm 1.0 kcal/mole and 43.6 \pm 1.8 kcal/mole for $Sb_2O_3(o)$.

The heats of formation (298°K) for the antimony sesquioxide are as follows: $\Delta H_{f(c)}^{O} = -164,430$ cal/mole (cubic) and $\Delta H_{f(0)}^{O} = -167,210$ cal/mole (orthorhombic), and for free energies, $\Delta G_{c}^{O} = -149,690$ cal/mole and $\Delta G_{o}^{O} = -147,900$ cal/mole. (51)

The specific heat from 273-929°K for $8b_20_3$ is reported by Maier⁽⁵¹⁾ as follows:

$$Cp_{Sb203} = 19.1 + 0.0171T$$
 $T = K$ (3.9)

and for Sb₂O₄ over undefined temperature range,

$$C_{P_{Sb204}} = 22.6 + 0.0162T$$
 $T = K$ (3.10)

It is assumed that $\mathbf{C_p}$ for cubic and orthorhombic forms of $\mathbf{Sb_2O_3}$ are identical.

In a vibrational and thermal study of the antimony oxides, Cody, DiCarlo and Darlington (52) found that both $\mathrm{Sb}_2\mathrm{O}_3(c)$ and $\mathrm{Sb}_2\mathrm{O}_3(o)$ when heated to $1000\,^\circ\mathrm{C}$ under nitrogen lose weight by a two-stage process: sublimation of the solid followed by volatilization of the melt. Examination of the thermograms presented by those authors indicate that $\mathrm{Sb}_2\mathrm{O}_3(c)$ loses weight over a smaller temperature interval than $\mathrm{Sb}_2\mathrm{O}_3(o)$ and that volatilization of $\mathrm{Sb}_2\mathrm{O}_3(c)$ is complete at a lower temperature than $\mathrm{Sb}_2\mathrm{O}_3(o)$.

The oxidation studies of the trioxide dimorphs by Cody, et al. $^{(52)}$ addressed some controversy generated in earlier thermogravimetric analysis (TGA) and differential thermal analysis (DTA) studies by Agrawal, et al. $^{(53)}$ In that study, TGA and DTA data for cubic Sb_2O_3 in inert and oxidizing atmospheres was reported. They claimed a TGA weight gain under oxidizing conditions of 4.5% for $Sb_2O_3(c)$ which is slightly less than the 5.49% expected for the following reaction if no sublimation of the starting material occurs:

$$28b_2o_3 + o_2 + 28b_2o_4 \tag{3.11}$$

Agrawal, et al., $^{(53)}$ data for cubic ${\rm Sb}_2{\rm O}_3$ is similar to that of Cody, et al., for orthorhombic ${\rm Sb}_2{\rm O}_3$ and may be

rationalized if dimorphic identification by Agrawal, et al., was incorrect.

The oxidation at 490°C of orthorhombic Sb₂O₃ to orthorhombic Sb204 (Sb203 . Sb205) was found to be topotactic by Gopalakrishnan and Manohar(54) who used x-ray techniques to determine structure of the reaction product. In a topotactic reaction there is a threedimensional orientational relationship between the original and final lattice. The same investigators calculated shifts in individual atom positions during oxidation is no more than 0.6Å. The oxidation was thought to be the result of oxygen moving along empty channels in the $Sb_2o_3(o)$ structure to the reaction site. Then an oxygen atom bridges two antimony atoms of neighboring chains. Calculations based on unit cell dimensions indicated that there is a 7.4% volume contraction when $Sb_2O_3(o)$ is oxidized to Sb₂O₄. Additional experiments indicated that the reduction of Sb_2O_4 to $Sb_2O_3(o)$ under hydrogen (430°C, 1 atm., 4 hrs.) is also topotactic. Sb₂O₃(o) was formed, not $Sb_2O_3(c)$, even though $Sb_2O_3(c)$ is stable at the reaction temperature.

Those authors $^{(55)}$ also determined in poly and single crystal $\mathrm{Sb}_2\mathrm{O}_3(o)$ oxidation studies that the rate law is parabolic suggesting an oxygen diffusion mechanism as the rate controlling step. Using a polarized light method for single crystals, they determined the activation energy for

two crystallographic directions. The activation energies were found to vary from 36 to 55.5 kcal/mole depending upon the crystallographic axis observed. The fastest reaction occurred through openings between chains of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$. The authors concluded that since the openings are not sufficiently large for simple free oxygen diffusion, the mechanism probably consisted of making, breaking and remaking bonds between oxygen and antimony atoms. A similar oxidative study of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ has not been reported in the literature.

Harmer and Pantano⁽⁸⁾, in a study of the synergistic interaction of Sb_2O_3 additions to the solid lubricant MoS_2 , obtained TGA data for those materials. Those investigators noted a 21.7% weight loss for Sb_2O_3 . They did not identify the dimorph evaluated nor the TGA heating rate. A proposed reaction to rationalize the weight loss was given as follows:

$$45b_2o_3 + \frac{5}{2}o_2 + 35b_2o_4 + 5b_2o_5 \tag{3.12}$$

followed by

$$5b_2o_5 + 25b + \frac{5}{2}o_2 +$$
 (3.13)

The hypothesis was justified by the handbook decomposition temperature (28) of Sb_2O_5 reported to be

400°C, near the 410°C inflection temperature observed in the thermogram, and the calculated weight loss is 20.9% in good agreement with the observed value. However, decomposition temperatures reported by Golunski, et al., (56) in more recent experiments where Sb₂O₅ gradually loses O₂ at temperatures above 400°C and decomposes as follows at higher temperatures:

$$Sb_2O_5(c)$$
 $650-700^0C$ + $Sb_2O_4(o)$ (stable to 1000^\circC) (3.14)

This decomposition reaction and temperature do not support the overall mechanism proposed by Harmer and Pantano. (8)

In the subsequent comprehensive study, Cody, et al., $^{(52)}$ obtained additional oxidative thermograms. In those studies, it was found that when the temperature of either $\mathrm{Sb}_2\mathrm{O}_3(c)$ or $\mathrm{Sb}_2\mathrm{O}_3(o)$ is increased rapidly $(20^{\circ}\mathrm{C/min})$ in air, the product is orthorhombic $\mathrm{Sb}_2\mathrm{O}_4$. However, $\mathrm{Sb}_2\mathrm{O}_3(c)$ experiences a 21% weight loss while $\mathrm{Sb}_2\mathrm{O}_3(o)$ gains 0.8% weight. They claimed that the competing processes of sublimation and reaction of $\mathrm{Sb}_2\mathrm{O}_3(c)$ and $\mathrm{Sb}_2\mathrm{O}_3(o)$ are responsible for the weight loss and gain, respectively.

Thus, it is seen that if Harmer and Pantano⁽⁸⁾ evaluated $Sb_2O_3(c)$ and if their TGA heating rate was sufficiently less than that of Cody, et al.,⁽⁵²⁾ then their data are in agreement with that of Cody, et al., but are in disagreement with that of Agrawal, et al.,⁽⁵³⁾. Comparison

of thermograms suggests that Agrawal, et al., may have oxidized ${\rm Sb}_2{\rm O}_3({\rm o})$ rather than ${\rm Sb}_2{\rm O}_3({\rm c})$ as noted earlier. These contradictory data suggest that the Agrawal, et al., DTA on ${\rm Sb}_2{\rm O}_3({\rm c})$ is suspect.

Golunski, et al., (56) resolved some of the issues generated in earlier studies in their comprehensive study of the thermal stability of the oxides of antimony. They found that Sb₂O₃(o) produced a large DTA endotherm under flowing nitrogen with onset at 628°C and peak at 643 ± 2°C. By TGA, weight loss was not significant until temperatures above 625°C under nitrogen were reached. Infrared and x-ray analysis of the cooled DTA product indicated no change. Therefore, it was concluded that in an inert atmosphere, orthorhombic Sb₂O₃ begins to melt at 628°C after which the liquid formed volatilizes.

In flowing air, DTA for $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ exhibited a large irreversible exotherm with a peak at $463^{\circ}\mathrm{C}$ with TGA indicating large changes in output depending upon experimental conditions. Infrared and x-ray analysis confirmed that $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ is converted to $\mathrm{Sb}_2\mathrm{O}_4$ in the temperature range of $500\text{--}700^{\circ}\mathrm{C}$. $\mathrm{Sb}_2\mathrm{O}_4$ was found to be stable to a temperature of $1105^{\circ}\mathrm{C}$.

In similar experiments with $\mathrm{Sb}_2\mathrm{O}_3(c)$, Golunski, et altound in DTA analysis that the dimorph in flowing nitrogen exhibited two peaks. The first, which was irreversible with onset at 629°C and peak at 639°C, was attributed to

the transformation to $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$. This was confirmed by infrared analysis of the product. The second peak, reversible, with an onset at 643 \pm 2°C and peaked at 651 \pm 2°C was related to as the melting and immediate progressive vaporization of molten $\mathrm{Sb}_2\mathrm{O}_3$.

For DTA experiments, the heating rate was 10° C/min. For nitrogen or air, the flow rate was 500 cm³/min. In TGA experiments, the heating rate was 15° C/min with a flow rate of either air or nitrogen of 200 cm³/min.

The data of Golunski, et al., (56) confirm the work of Cody, et al., (52) regarding the inaccuracies in the data presented by Agrawal, et al. (53) DTA for the oxidation of Sb2O3(c). The data also offer an alternative to the mechanism proposed by Harmer and Pantano(8) for oxidation of Sb2O3 generated in their TGA study. That is, the Sb2O3 (unknown structure) thermally analyzed in air by Harmer and Pantano was, most probably, oxidized while a portion of the sample volatilized simultaneously thus explaining their results. DTA and TGA results are obviously dependent upon the dimorph used, particle size and distribution, heating rate and atmospheric conditions which offer an explanation for the varied results generated experimentally and the hypotheses proposed to rationalize the data.

It is noted that in addition to orthorhombic $\alpha + Sb_2 U_4$, formed in the above oxidation reactions, another polymorph,

 $\beta-5b_20_4$ (monoclinic), exists as reported by Rogers and Skapski. (57) The structure was determined using x-ray methods. Since it is claimed that $\alpha-5b_20_4$ forms in air or oxygen, Rogers and Skapski suggested the following mechanism for the transformation:

$$2\alpha-Sb_2O_4(s) + Sb_4O_6(vapor) + O_2 + 2\beta-Sb_2O_4(s)$$
 (3.15)

In more recent studies by Cody, et al., (52) it is suggested that the above mechanism may be erroneous, since β -Sb₂O₄ was not produced when α -Sb₂O₄ was exposed to flowing oxygen at 1130 - 1180°C. Evidently, β -Sb₂O₄ is a stable high pressure form of Sb₂O₄, since β -Sb₂O₄ was formed in a few minutes at 960°C in sealed capillaries (~5 atm) by the latter investigators. The mechanism of transformation is still uncertain. Temperature and pressure for the transformation are yet to be adequately defined.

Analytical methods employed in the above and other studies of antimony oxides which have proven useful include x-ray, (44,52) infrared, (48,52) Raman, (52) Mossbauer, (44) XPS, (58,59) and Auger (59,60) spectroscopy.

PROTECTIVE INTERACTION OF Sb203 ADDITION TO MoS2

The first description of the use of ${\rm Sb}_2{\rm O}_3$ in a ${\rm MoS}_2$ solid lubricant system was made in a 1966 patent by Murphy. (3) He claimed that solid film epoxy-phenolic-

bonded solid lubricants containing both $\mathrm{Sb}_2\mathrm{O}_3$ and MoS_2 with other inorganic or organic components exhibited Falex test mathine wear lives of 500 minutes, and time to failure for coated 1020 steel panels under 20% salt spray of 120 hours. In his formulation, the weight percentage of $\mathrm{Sb}_2\mathrm{O}_3$ was 6.2% while MoS_2 was 17.4%.

Since the granting of Murphy's patent, there have been several studies directed toward developing an optimum ${\rm Sb_2O_3\text{-}MoS_2} \ \ {\rm formulation} \ \ {\rm with} \ \ {\rm only} \ \ {\rm minor} \ \ {\rm effort} \ \ {\rm devoted} \ \ {\rm to} \ \ {\rm examining} \ \ {\rm mechanisms} \ \ {\rm repsonsible} \ \ {\rm for} \ \ {\rm the} \ \ {\rm protective} \ \ {\rm role} \ \ {\rm of} \ \ \ {\rm Sb_2O_3} \ \ \ \ \ {\rm when} \ \ \ {\rm added} \ \ \ {\rm to} \ \ {\rm MoS_2}.$

Following several exploratory efforts (4,5) to develop optimum resin-bonded film performance, the first study to offer an interaction rationale for the Sb203-MoS2 system was given by Lavik, Hubbell and McConnell. (7) In 1975, they offered the hypothesis that an eutectic of Sb203 and MoO3 forms on a lubricating compact of Sb203 and MoS2. During sliding interfacial oxidation at hot asperity sites would provide materials for eutectic formation so that additional enhanced transfer films formation occurs. Those authors evaluated resin-bonded MoS2/Sb203 lubricants, 'n a dual rub-shoe test, bonded films of MoS2/Sb203 using the Falex method, and compacts of MoS2/Sb203 in a pellet test. The addition of Sb203 improved the wear life of MoS2 in the resin-bonded materials. Significantly, when the effects of the resin binder were eliminated in a pellet

study, wear life was again superior for MoS_2 compacts containing 45 volume percent Sb_2O_3 compared with MoS_2 compacts without Sb_2O_3 .

The authors did not offer any evidence to support the mechanism proposed. That is, no evidence for the existence or formation of the proposed eutectic was given, rather only test data indicating superiority of the MoS_2/Sb_2O_3 lubricant as cited above was provided.

They also noted that improvement in coefficient of friction was found for the compounded lubricant compared with noncompounded ${\tt MoS}_2$.

A subsequent study of the system was conducted by Harmer and Pantano⁽⁸⁾ who reviewed the work of Hopkins and Campbell⁽⁶¹⁾ noting that the study was inconclusive regarding the properties of the MoS₂-Sb₂O₃ solid lubricant system. Harmer 'd Pantano stated that the inconsistencies in data reported for the MoS₂-Sb₂O₃ system might be related to the use of polyimide as a resin binder as compared with MoS₂-Sb₂O₃ bonded with a phenolic-epoxy. Close examination of the data presented indicates that some test were run in dry atmosphere while or 'rs were conducted in relatively high humidity conditions. The effect of water on the polyimide available at that time is well-known; ^(62,63) thus, the data can be remained since the polyimide composite was far superior to the phenolic-epoxy and all others in a nitrogen pellet test, yet inferior to the

phenolic-epoxy when run in ambient air journal bearing tests. Therefore, it appears that the Hopkins and Campbell (61) data can be satisfactorily rationalized based solely on consideration of the deleterious effects of water on the properties of polyimides.

Harmer and Pantano (8) conducted a preliminary study directed towards the elucidation of the interaction of MoS2 and Sb_2O_3 as referred to earlier. They employed thermogravimetric methods (TGA), x-ray diffraction, and Auger analyses of compacted and burnished materials to generate some basic data on the interaction. Among their conclusions were that: (i) distinct evidence of chemical interaction ${\rm Sb_2O_3/MoS_2}$ was found in TGA (air) studies; (ii) MoS_2 was oxidized to MoO_3 and Sb_2O_3 to Sb_2O_4 as shown by x-ray studies; (iii) Sb203 or its oxidation product, Sb204, did promote sintering of MoO3; (iv) MoO3 showed little solubility in $\mathrm{Sb}_2\mathrm{O}_4$, and conversely, $\mathrm{Sb}_2\mathrm{O}_4$ showed little solubility in MoO_3 ; (v) Auger studies indicated that Sb_2O_3 containing treated pellet surfaces were significantly different from those of pure MoS2; and (vi) Mo concentrations near pellet surfaces of MoS2/Sb203 were anomalously low. Rationale for these results was not presented.

Harmer and Pantano indicated that additional efforts should be expended to determine if (i) the microstructures of the solid lubricant transfer film changes with ${\rm Sb}_2{\rm O}_3$

addition; (ii) segregation of Sb_2O_3 or Sb_2O_4 occurs at the sliding interface; (iii) there is a role for water vapor; and (iv) changes in the surface chemistry affect the lubricant properties of MoS_2/Sb_2O_3 films, especially the role of sulfur.

Although Harmer and Pantano⁽⁸⁾ claimed to find evidence of interaction and thought that eutectic formation might play a role, no direct evidence of the formation of a eutectic was offered. Because of lack of funding, their effort was, unfortunately, discontinued.

A subsequent effort by Nosov⁽⁹⁾ in 1978 reported upon the lubricating properties of molybdenum disulfide with additives of antimony oxide and antimony sulfide. Nosov conducted an empirical study of MoS₂ coatings containing various concentrations of Sb₂O₃ or Sb₂S₃. Those tests determined the time required to abrade coating in both sliding and rolling contact. Nosov concluded that the antimony compounds showed high tribochemical activity and improved the lubricating properties or MoS₂ base coatings. He further stated without evidence that the formation of an eutectic in the contact zone was not very likely under standard test conditions because of temperatures as proposed by Lavik, et al.⁽⁷⁾ Nosov also claimed without direct evidence that antimony trioxide increases the oxidation resistance of MoS₂ coatings.

The most recent work in which MoS_2/Sb_2O_3 formulated solid lubricants were used is that of Gardos and McConnell. (1) In the study potential high-temperature solid lubricants were evaluated in resin-bonded dual rub-shoe test configuration. Sb_2O_3/MoS_2 lubricant was employed as the standard. The reference shoe composition was 40 weight percent MoS_2 , 30 weight percent Sb_2O_3 , and 30 weight percent polyimide.

They compared various other solid lubricants to the above reference formulation. The results indicated that only one of the studied formulations was marginally superior to the standard MoS₂-Sb₂O₃ formulation. Gardos and McConnell's concluding comment regarding the protective mechanism of Sb₂O₃ additions to MoS₂ was that, based upon all the evidence to date, Sb₂O₃ has an oxygen scavenging role, i.e., in mixtures of MoS₂ and Sb₂O₃, the Sb₂O₃ will be preferentially oxidized. No experimental evidence was offered to support their conclusion.

The above summarizes the significant work concerning the apparent synergism of ${\rm Sb_2O_3}$ additions to ${\rm MoS_2}$ which provide improved tribological performance.

In review of the hypotheses, it is plausible that an eutectic of $\mathrm{Sb}_2\mathrm{O}_3$ and MoO_3 might be formed at the sliding interface and in some manner improve tribological properties, since eutectics of $\mathrm{Sb}_2\mathrm{O}_3\text{-MoO}_3$ are known. $^{(64)}$ The second hypothesis, proposing sacrificial oxidation of

 ${\rm Sb}_2{\rm O}_3$ to ${\rm Sb}_2{\rm O}_4$ or higher oxides thus preventing oxidation of ${\rm MoS}_2$, lacks plausibility since earlier workers had found that ${\rm Sb}_2{\rm O}_3$ additions were beneficial even when evaluated in vacuum. Thirdly, hypotheses involving an assumed role for ${\rm Sb}_2{\rm O}_3$ in retardation of ${\rm MoS}_2$ oxidation have not been sufficiently defined to critically analyze without additional data. The eutectic and oxidation retardation hypotheses will be discussed in greater detail.

The eutectic hypothesis proposes that is might be possible for an eutectic of oxides to promote attachment of a MoS₂ transfer film to a sliding surface. Although consideration of the possible role of eutectic formation and/or oxygen scavenging by Sb₂O₃ was given by some investigators, (1) none of the workers sought or found the Sb₂O₃-MoO₃ phase diagram in the literature as reported by Parmentier, Courtois and Gleitzer in 1974. (64)

Coincidentally, the empirically optimized formulations, found by Lavik, et al., (7) and Gardos and McConnell, (1) are very close in composition to that of the highest melting point eutectic composition found by Parmentier, et al., which would be derived from a compact containing the largest fraction of MoS₂.

If the eutectic performs a role, then the selected formulations may be rationalized as resulting from the need to form the compact with as much ${\rm MoS}_2$ as possible since it is obviously the primary lubricant along with the need to

form a low-temperature eutectic for improved tribological properties. Such a point is near the experimental formulations (1,7) which appear to offer the best combination of properties. The coincidence of a eutectic point with typical formulations does not provide direct evidence for the role of a eutectic, but it is obvious that the possibility exists from a study of the phase diagram.

However, if one considers that investigators $^{(65)}$ have estimated a typical MoS_2 film is $1-2 \times 10^{-6}\mathrm{m}$, and since the distance between molybdenum layers in MoS_2 is $6.16 \times 10^{-10}\mathrm{m}$, $^{(66)}$ then under typical sliding conditions more than 3,000 layers of MoS_2 might be between the substrate and sliding interface. The thickness of MoS_2 weighs against attachment of $\mathrm{h...S}_2$ to a substrate by an oxide as being the critical mechanism involved in improvement of tribological properties when such bulk additions are required. Because there are nearly 3,000 layers of MoS_2 between the site of the proposed effect and the sliding zone, it would appear that an alternative mechanism involving the entire solid lubricant film structure would be more plausible.

One possible mechanism for $\mathrm{Sb}_2\mathrm{O}_3$ retarding MoS_2 oxidation could be similar to the mechanism in which it is employed as an antitoxidant in plastics and other solids. (67) However, it is known that the presence of a halogen is required for $\mathrm{Sb}_2\mathrm{O}_3$ to act effectively as an

antioxidant involving the formation of SbCl3 where Cl is an example of a halogen as follows:

$$Sb_2O_3 + 2HC1 + 2SbOC1 + H_2O$$
 (3.16)

followed by

$$SbOC1 + 2HC1 + SbC1_3 + H_2O$$
 (3.17)

 ${
m SbGl}_3$ acts as both a dehydrating agent and a free radical chain terminator. In the ${
m MoS}_2{
m -}{
m Sb}_2{
m O}_3$ formulations evaluated previously, none contained halogens. Alternatively, if the mechanism is assumed to be one where ${
m Sb}_2{
m O}_3$ acts to block diffusion of oxygen into the ${
m MoS}_2$ matrix, thereby retarding oxidation, then many other materials would be effective additives which previous investigators did not claim.

retardant, (68) although the mechanism is evidently not similar to Sb₂O₃. In polyvinyl chloride, MoO₃ acts to isomerize trans conjugated polyene structures produced on the loss of hydrogen chloride which is the first step in the pyrolysis of polyvinyl chloride. (69) The cis isomer produced in the presence of MoO₃ as a Lewis acid is more thermally stable. Therefore, the flame retardant mechanism of MoO₃ is totally dissimilar to that of Sb₂O₃ which is also used in polyvinyl chloride. The mechanisms by which

 ${\rm Sb_2O_3}$ and ${\rm MoO_3}$ act as flame retardants for other materials do not appear to be possible in ${\rm MoS_2-Sb_2O_3}$ compacts.

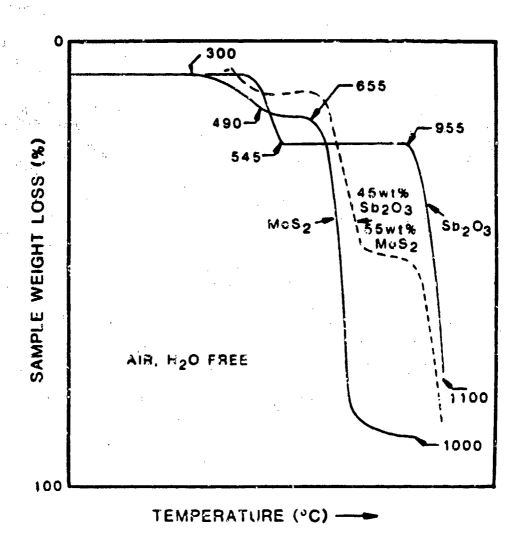
CHAPTER IV

ROLE OF OXIDATION OF ANTIMONY OXIDE

An understanding of Sb₂O₃ oxidation is required for elucidation of its beneficial role in MoS2 compacts, since previous investigators have suggested that such ${\rm Sb}_2{\rm O}_3$ oxidation may be critical. For example, Gardos and McConnell(1) proposed, without experimental data to support the concept, that Sb_2O_3 is preferentially oxidized rather than MoS2. Thus, the sacrificial oxidation of Sb203 would reduce the rate of oxidation of MoS_2 to MoO_3 and to molybdenum sulfate compounds, which would result in reduced wear while maintaining low friction. With respect to this hypothesis, Harmer and Pantano (8) in thermal gravimetric apparatus (TGA) studies found that MoS2 started to oxidize at 300°C with oxidation to MoO3 being completed by 490°C followed by loss of MoO3 by sublimation. They also found that Sb₂O₃ initially lost weight at 410°C followed by more rapid weight loss until 545°C. Then the weight remained constant until 955°C. Harmer and Pantano interpreted these data as oxidation of Sb203 to Sb204 and rationalized the unexpected weight loss as resulting from a reaction involving the formation of Sb205, which then decomposed to Sb sublimate and O2. They did not consider sublimation of

Sb203, which was found to occur simultaneously with oxidation in similar experiments by Cody, et al. (52) The net effect of oxidation and sublimation of Sb203(c) resulted in a 21 percent weight loss nearly identical with the Harmer and Pantano results.

Harmer and Pantano also obtained TGA data for mixed MoS₂ (75 weight percent) and Sb₂O₃ (25 weight percent). In dry air, they found a slight increase in weight at 348°C followed by weight loss until 468°C. Then the weight remained stable until a slight increase at 550°C followed hy very dramatic weight loss from about 650° to 863°C. These data, based upon Cody, et al., conclusions can be raionalized as a combination of KoS2 and Sb202 oxidation, . Sb_2O_3 and MoO_3 sublimation, followed by eventual sublimation of Sb₂O₄, without proposing a mechanism wherein Sb₂O₃ retards oxidation of MoS₂. Especially significant in Harmer and Pantano's analysis was a conclusion that since the weight loss at 480°C was only 9.5 weight percent rather than nearly 14 weight percent predicted by theory, then some role for Sb_2O_3 in retarding MoS_2 oxidation must be assigned. Again, however, simultaneous oxidation and sublimation of Sb_2O_3 was not considered. Harmer and Pantano also analyzed a 45 weight percent Sb203 55 weight percent MoS2 mixture. TGA results for that mixture of Harmer and Pantano(8) are redrawn in Figure 1 where those data are compared with oxidation of MoS2 and Sb203. Weight



Pigure 1. Compilation of Harmer and Pantano's Theimal Analysis Date. (8)

losses and temperatures of inflections were slightly different from those found for the 25 weight percent $\mathrm{Sb}_2\mathrm{O}_3$ -75 weight percent MoS_2 data, however, differences in the analyses may be rationalized if sublimation of $\mathrm{Sb}_2\mathrm{O}_3$ and its effect on reaction rates are considered.

Therefore, because of the evident need to examine the role of antimony oxide oxidation as a beneficial mechanism of $\mathrm{Sb}_2\mathrm{O}_3$ additions to MoS_2 , and the fact that isothermal kinetic data for the oxidation of $\mathrm{Sb}_2\mathrm{O}_3(c)$ have not been reported, which might be significant in such a protective role, a study of the oxidation of $\mathrm{Sb}_2\mathrm{O}_3$ and its possible influence on lubricant compact performance was undertaken.

ISOTHERMAL OXIDATION OF Sb203

Isothermal kinetic studies were undertaken using various particle sizes of the dimorphs of ${\rm Sb}_2{\rm O}_3$. Each dimorph and mixtures were oxidized. The method was to employ a closed glass reaction system that could be used to maintain oxygen at nearly constant pressure (140 torr) and that would permit measurement of the volume of ${\rm O}_2$ consumed by means of an ${\rm O}_2$ -filled gas buret as given in figure 2. A Stone differential thermal analysis (DTA) Model F-1DF furnace was used to maintain a constant reaction tube temperature, which was measured using a chromel-alumel thermocouple accurate to $\pm 0.4\%$. This system is similar to the one used by Gopalakrishnan and Manchar in their

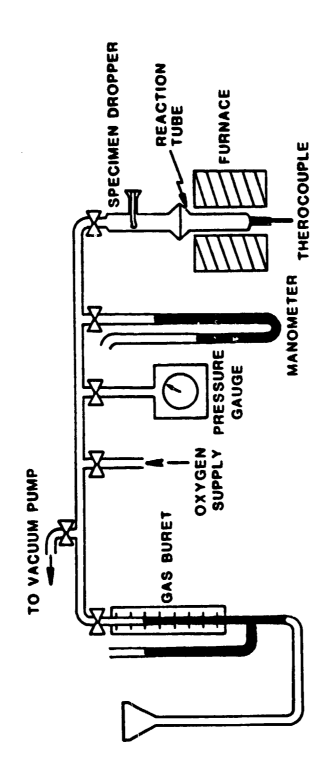


Figure 2. Isothermal Oxidation Apparatus.

oxidation studies of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$. The experimental procedure is given in Appendix A.

Sb203 for reaction studies was prepared from 99.999% pure (metals basis) Sb203 (Alfa, Danvers, MA). Materials, were placed in a closed Pt cylinder which was then placed in a quartz tube. After evacuating the quartz tube three times to 10⁻⁶ atm and refilling with 99.9995 percent pure argon (Matheson, East Rutherford, NJ), the tube was filled to 0.2 atm with argon after which the tube was sealed. The quartz tube was then placed into a muffle furnace adjacent to an indicating thermocouple. Desired muffle furnace temperatures were obtained within an hour with set point temperatures at about 520-540°C for Sb203(c) and about 620-640°C for Sb203(o) production. Oven temperatures were proportionally controlled throughout the heat treatment. Photographs of typical Sb203 products obtained by scanning electron microscopy are presented in Figures 3 and 4.

The infrared spectrometric method used by Gopalakrishnan and Manohar $^{(48)}$ as developed by Borger and Krogh-Moe $^{(70)}$ was used initially to analyze both reactants and products. The accuracy $^{(48)}$ of the method based upon analysis of dimorph mixtures indicated that accuracy was limited to $\pm 5\%$.

Another method used for analysis was Raman spectrometry as reported by Cody, et al.(52) The Raman spectra for $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$, $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$, and $\mathrm{Sb}_2\mathrm{O}_4$ consists of



Figure 3. SEM Photograph of Sb₂0₃(c).

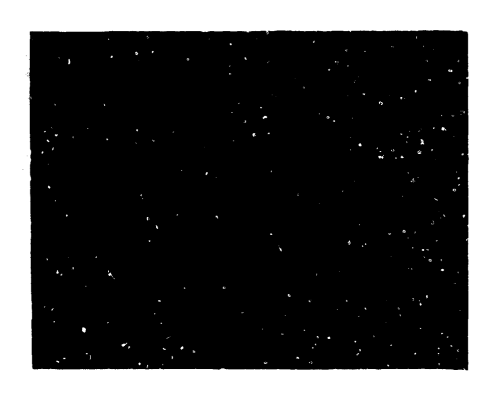


Figure 4. SEM Photograph of Sb₂0₃(o).

sharp, well-separated peaks useful for qualitative purposes. Analyses of known mixtures suggested that packing variations of powders in Raman tubes of different particle sizes adversely affected the accuracy of analyses. It was also observed that Raman intensities decreased markedly as $Sb_2O_3(o)$ crystals were ground into fine powders. These results are in agreement with those of Ferraro, et al. (71) Therefore, analysis performed using either technique was employed on a semi-quantitative basis only.

ISOTHERMAL OXIDATION OF Sb203(o)

The isothermal oxidation of Sb₂O₃(o) was reported by Gopalakrishnan and Manohar⁽⁵⁵⁾ to take place at increasing xates of reaction from 470° to 520°C. Their reaction rates for very large crystal size Sb₂O₂(o) (2 x ·1 x ·02 mm) could not be reproduced for crystals of about that size. It was found that similar crystals did not oxidize at 470°C after 50 minutes. They report over 65 percent theoretical completion after 50 minutes at 470°C under oxygen at 140 torr. In this work it was found that such large crystals could only be oxidized to 60 percent theoretical completion if the reaction temperature was increased, e.g., a temperature of 533°C under oxygen at 163 torr for one hour.

It was then decided to grind $\mathrm{Sb}_2\mathrm{O}_3(o)$ to a much smaller particle size to determine if the oxidation rate would be increased for samples with more surface area.

Therefore, a sample of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ was ground mechanically on a mechanical amalgamator (Wig-L-Bug, Cresent Dental, Chicago, IL) for 16 minutes. Samples were analyzed by both infrared and Raman spectrometric techniques which always indicated that $\mathrm{Sb}_2\mathrm{O}_4$ had not been formed during the grinding process.

The 16-minute ground $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ was then oxidized isothermally in the kinetics rig at temperatures ranging from $453\,^\circ\mathrm{C}$ to $521\,^\circ\mathrm{C}$. These data are presented in Appendix B and Figure 5. In these experiments it was found that the theoretical limit of oxygen consumption could only be approached at higher temperatures. At such reaction temperatures, a light coating on the interior of the reaction tube identified as $\mathrm{Sb}_2\mathrm{O}_4$ was found at increasing height above the reactant at higher reaction temperatures, but $\mathrm{Sb}_2\mathrm{O}_4$ was never found at more than about one cm above the top of the solid sample.

Repeat of isothermal oxidation of $5b_20_3(o)$ at $521^{\circ}C$ indicated good repeatability for oxidation conducted in the glassware system as presented in Figure 6.

Examination of the reaction data indicates that after an initial induction time, which shortened as the reaction temperature increased, the oxidation of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ appears to be directly proportional to the reaction time. After some time, the reaction slows down and appears to become proportional to the square root of time, a parabolic rate,

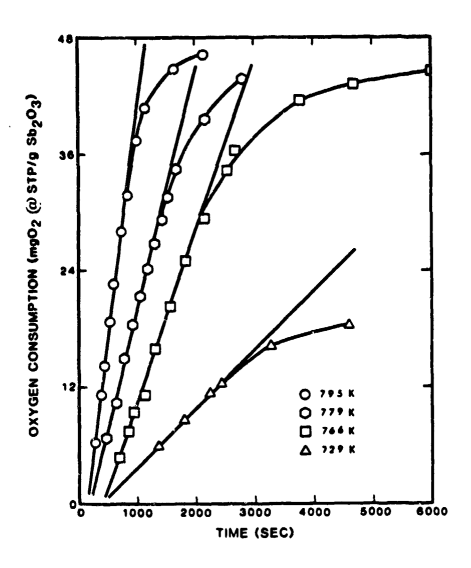


Figure 5. Isothermal Oxidation of $Sb_2o_3(o)$.

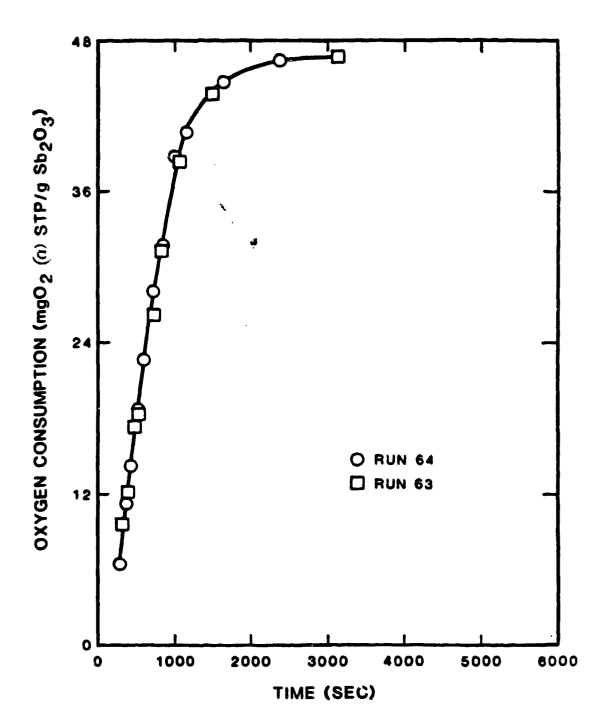


Figure 6. Repeatability of Sb203(o) Oxidation.

or even slower for later segments of the reaction. Oxygen consumption as a function of the square root of time is presented in Figure 7.

Trofinov and co-workers (72) have provided a basis for such a change of reaction rates during the reaction. They found in an x-ray heated cell experiment that fine powdered Sb203(0) oxidized at lower temperatures than did equivalent fine $Sb_2O_3(c)$. However, when mixtures $Sb_2O_3(o)$ and Sb₂O₃(c) were oxidized, Sb₂O₃(c) oxidized prior to Sb₂O₃(o). They rationalized those reactions by hypothesizing that oxidation is nucleated in a low energy process when orthorhombic Sb_2O_4 is formed coherently on orthorhombic Sb203(o) surface. Because Sb204 also provides such a surface, oxidation continued by sublimation of $Sb_2O_3(c)$, which has a higher equilibrium sublimation pressure than $Sb_2O_3(o)$, until the supply of $Sb_2O_3(c)$ is exhausted. Oxidation of covered Sb203(o) may continue but the process is controlled by a slow bulk diffusion process compared with gaseous diffusion of sublimed Sb406.

Thus, for the isothermal oxidation of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$, the process may be initiated by sublimation of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$, or more correctly $\mathrm{Sb}_4\mathrm{O}_6$, which oxidizes on the $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ surface. Eventually, all readily sublimable $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ is oxidized with remaining $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ covered with $\mathrm{Sb}_2\mathrm{O}_4$. At this point, oxidation can only occur by bulk diffusion. Then, at some point, the process terminates when the $\mathrm{Sb}_2\mathrm{O}_4$

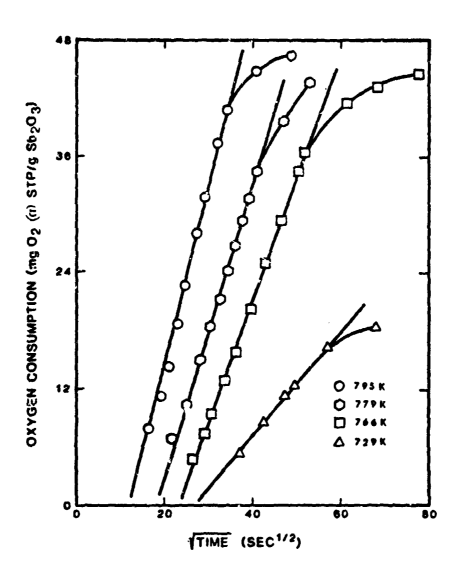


Figure 7. Parabolic Law Isothernic Oxidation of Sb203(0).

layer becomes too thick. In a plot of oxygen consumption as a function of reaction time, the initial section may be explained as sublimation of Sb_40_6 through some distance to a low energy nucleation site which continues linearly until coverage nears completion, at which time, a much slower bulk diffusion process becomes predominant. The rate of the bulk process not only slows rapidly, but any traces of sublimable reactant are oxidized quickly. If coverage is uniformly coherent and thick, then residual Sb_20_3 cannot be oxidized in a reasonable time and the reaction terminates.

In these experiments, if the temperature is very high, e.g., 521° C, then sublimation is dominant and the reaction proceeds to near theoretical limit. At lower temperatures, e.g., 456° C, the reaction forms Sb_2O_4 on $Sb_2O_3(o)$ surfaces more effectively, reducing the rate of sublimation so that the reaction becomes bulk diffusion controlled which results in termination of oxidation.

In contrast, Gopalarrishnan and Manohar $^{(55)}$ claim that reaction is completely oxygen diffusion controlled for the entire temperature range. Differences in distribution and sizes of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ crystals could provide the basis for the dissimilar hypotheses for the oxidation. However, the observations of Trofimov, et al., and the data presented here suggest that a sublimation controlled mechanism for oxidizing $\mathrm{Sb}_2\mathrm{O}_3$ to $\mathrm{Sb}_2\mathrm{O}_4$ is plausible, at least for a portion of the oxidation.

ISOTHERMAL OXIDATION OF Sb203(c)

 $Sb_2O_3(c)$ formed from Sb_2O_3 (Alfa) was produced during heat treating in smaller crystal sizes than $Sb_2O_3(o)$. The $Sb_2O_3(c)$, as found, was sieved through 20 micron onto 10 micron nickel mesh screus (0.336 m²/g BET surface area Micromerities, Morcross, GA).

Oxidation data are given in Figures 8 and 9. reaction rates are, like $Sb_2O_3(o)$, a function of comperature. Isothermal oxidation of Sb2O2(c) requires significantly higher temperatures than does SbyO3(o). The reaction rate is very slow at first, and quickly increases until some later time in the reaction. Depending upon the reaction temperature, the rate slows from an apparently linear region, like Sb₂O₃(o), to a parabolic rate, becoming even slower at latter stages until termination of the reaction. These rates generate a reaction curve having an elongated sigmoidal appearance. A plausible explanation is that cubic Sb_2O_3 does not have the low energy reaction $sites^{(72)}$ for an orthorhombic product; but as orthorhombic Sb₂O₄ is formed, it may provide the low energy nucleation site and become similar to $Sb_2O_3(o)$ in reaction characteristics.

 ${\rm Sb}_2{\rm O}_3(c)$ oxidized at 521°C in a used tube previously coated with the ${\rm Sb}_2{\rm O}_4$ product showed an increase in reaction rate up to about 50 percent of theoretical completion. This experiment suggests that the orthorhombic

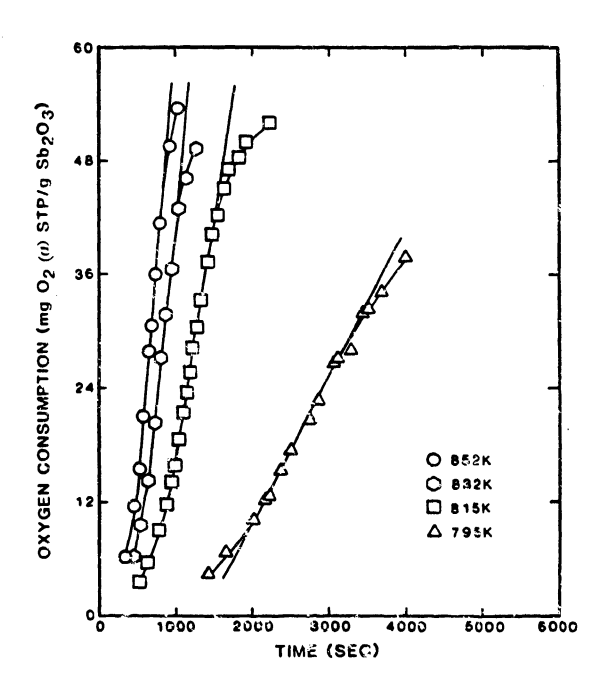


Figure 8. Isothermal Oxidation of $5b_20_3(c)$.

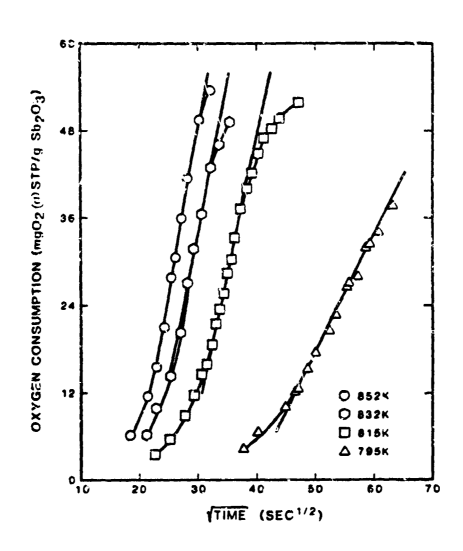


Figure 9. Parabolic Law Isothermal Oxidation of Sb203(c).

surface of ${\rm Sb_20_4}$ may be a low-energy nucleation site as suggested by Trofinov, et al. (72) These data are presented in Figure 10.

ISOTHERMAL OXIDATION OF MIXTURES OF $Sb_2O_3(c)$ AND $Sb_2O_3(c)$

50-50 weight percent mixtures of ground $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ and sieved, unground $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ were oxidized at temperatures from 492° to $530^\circ\mathrm{C}$. The results are presented in Figures 11 and 12. The reaction rates for the mixtures are intermediate between the pure dimorphs and as of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ and $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ appear to obey the same reaction mechanism. These data do not directly support the observation of Trofimov, et al. (72) that in mixtures of $\mathrm{Sb}_2\mathrm{O}_3$, the rate of oxidation is faster than would be predicted by summing of the rate for each dimorph. However, these oxidation rates employ larger $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ than that used in $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ oxidation rate determinations. These experiments were not repeated because after all other oxidation data was analyzed, an understanding of oxidation and its role in the beneficial mechanism of $\mathrm{Sb}_2\mathrm{O}_3$ in MoS_2 became clearer.

Use of sieved, unground $\mathrm{Sb}_2\mathrm{O}_3(c)$ may also provide an explanation for mixed oxidation data at 803 K compared with data at 793 K. Crossover of oxygen consumption near the end of reaction may result from significant differences in surface area of $\mathrm{Sb}_2\mathrm{O}_3(c)$ added to the mixture.

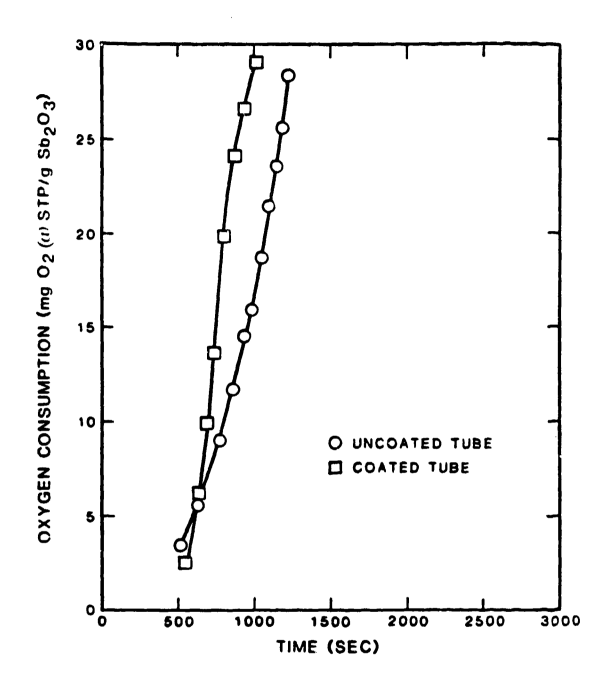


Figure 10. Oxidation of $Sb_2O_3(o)$ in New and Used Reaction Tubes.

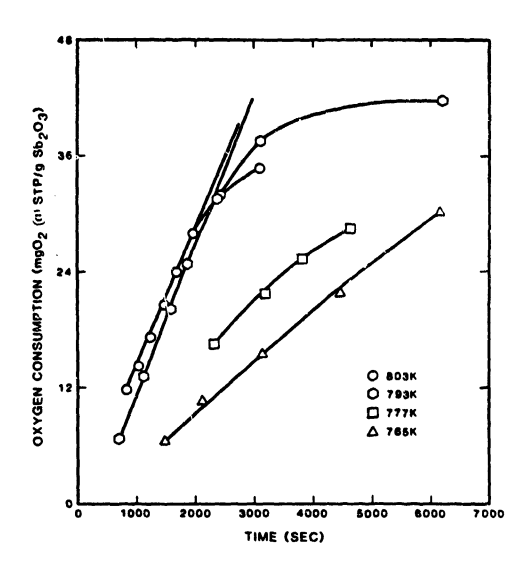


Figure 11. Oxidation of Mixed Dimorphic Sb203.

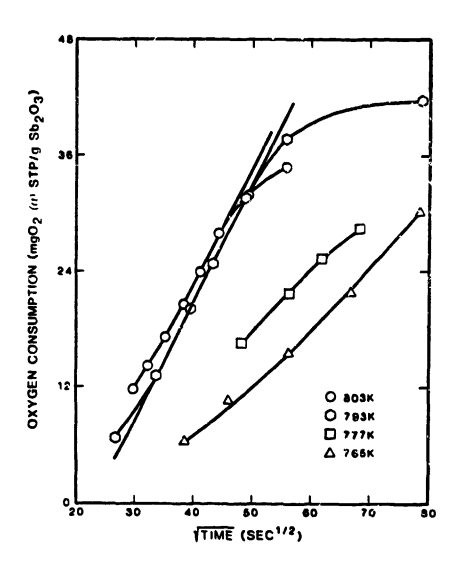


Figure 12. Parabolic Law Isothermal Oxidation of Mixed Dimorphic ${\rm Sb_2O_3}.$

EVALUATION OF ISOTHERMAL OXIDATION DATA

To determine the apparent activation energy for exidation of $\mathrm{Sb}_2\mathrm{O}_3(c)$, $\mathrm{Sb}_2\mathrm{O}_3(o)$ and mixed $\mathrm{Sb}_2\mathrm{O}_3$, the reaction rates were plotted as a function of the reciprocal of reaction temperature using the Arrhenius relationship as follows:

$$-E_{a}/RT$$

$$k - A e \qquad (4.1)$$

where

k = rate constant

A = pre-exponential or frequency factor

 E_a = apparent activation energy

R = gas constant

T - absolute temperature

By taking the logarithm of both sides,

$$\ln k = \ln A - \frac{E_a}{RT} \tag{4.2}$$

Then, when in k is plotted as a function of $\frac{1}{T}$, the slope is equal to $\frac{E_a}{R}$, while the intercept is in A. E_a is related to the energy required for the reaction to proceed. As calculated from data presented in Figures 13, 14 and 15, the mean E_a for $Sb_2O_3(c)$ is 48.0 ± 1.9 kcal/mole, 47.4 ± 10.0 kcal/mole for $Sb_2O_3(o)$, and 44.7 ± 0.8 kcal/mole for mixed Sb_2O_3 as given in Table 2. Least square lines were

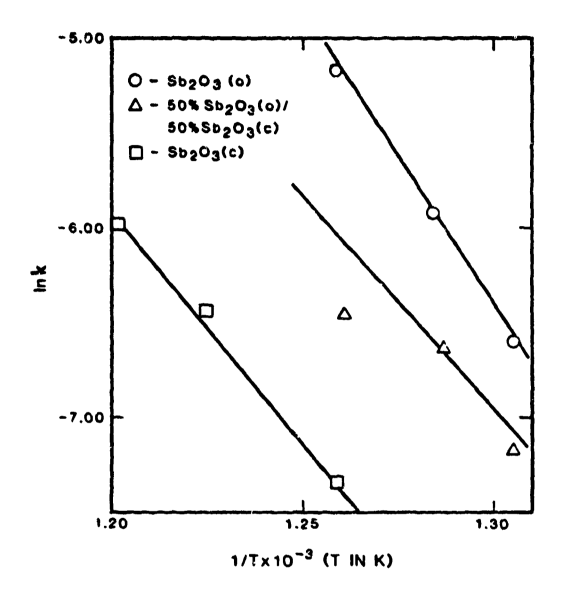


Figure 13. Arrhenius Plot for ${\rm Sb_2O_3}$ Oxidation at 10% Theoretical Reaction Completion.

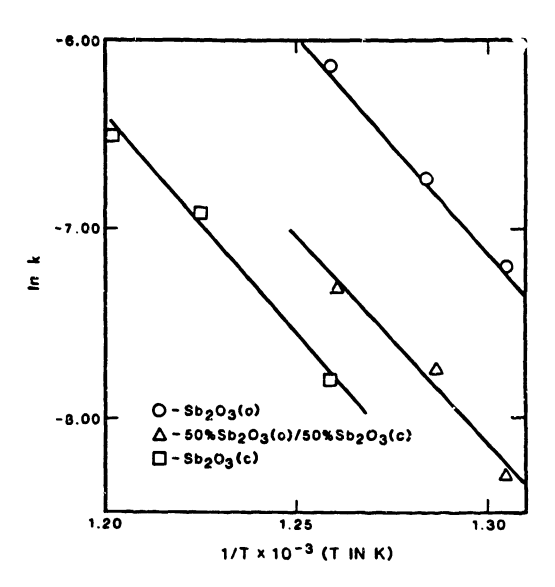


Figure 14. Arrhenius Plot for ${\rm Sb_2O_3}$ Oxidation at 30% Theoretical Reaction Completion.

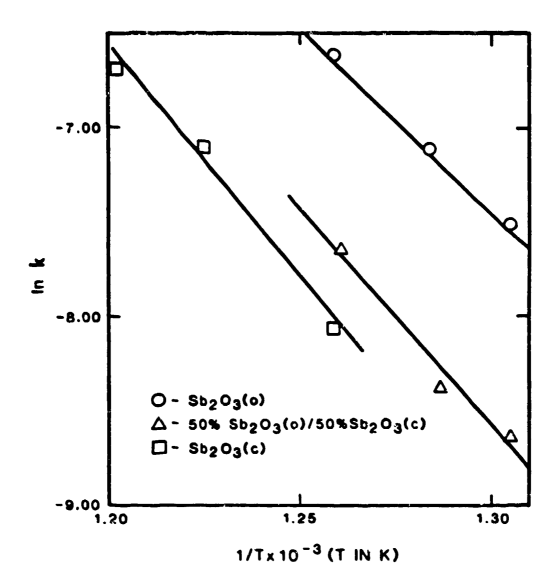


Figure 15. Arrhenius Plot for $\mathrm{Sb}_2\mathrm{O}_3$ Oxidation at 50% Theoretical Reaction Completion.

ACTIVATION ENERGIES (E,) AND PRE-EXPONENTIAL PACTORS (A) FOR THE ISOTHERMAL OXIDATION OF ${\rm Sb}_2{\rm O}_3({\rm c})$ Table 2.

							•	,		•					
Resctont		8b ₂ 0 ₃ (0)	(0)				8b 203(c)	(c)		-	Mixod	\$6203	Hized Sb203(e) - Sb203(c)	\$520	3
I Theoretical Completion	10 30	30	20	\$		30	30	50	\$	50 65 mean 10 30 50 65 mean 10 30 50 65 mean	10	36	2	3	i
E. (kcal/mole)	61.7 45.	45.8	.8 38.7 43.5 47.4 47.8 45.7 48.3 50.3 48.0 44.8 43.9 45.5 44.7	43.5	17.4	47.8	45.7	48.3	50.3	0.84	::	43.9	45.5	:	:
				•	#10					#:.					*0.
A (8 ⁻¹)	5.33	5.33 8.74 5.96 9.7 2.4 9.48 1.62 6.52 1.87 6.53 4.81 8.65 1.6	5.96	9.1	2.4	9.4	1 1.62	6.52		1 6.53	;	1 8.65	:	;	
	31014 11C	#10	09 x107 x108 x1010 x109 x109 x1010 x101 x109 x108 x109	# 10	11010	#109	*10°	*109	×101	\$01x C	*10°	# 0 T K	601×		*109

constructed for the data, except high temperature data for 10 percent oxidation of mixed Sb_2O_3 for which the oxidation appears to be unreasonable. The means are for apparent activation energies over the range of 10 to 50 or 65 percent reaction completion and standard deviations are for those data. Values of , the pre-exponential constant, are typically 10^9-10^{-1}

The computed values of E_a are very close to those of Jungermann and Plieth⁽⁵⁰⁾ who found that the enthalpy of sublimation (ΔH_B) of $Sb_2O_3(c)$ is 43.8 \pm 1.0 kcal/mole and 43.6 \pm 1.8 kcal/mole for $Sb_2O_3(o)$. From the similarity of E_a and ΔH_B , and the observation of product on reaction tube walls, one may hypothesize that under these psuedo-isobaric conditions, sublimation is the rate controlling step for the oxidation of the antimony trioxides. However, the role of oxygen was not determined, and the adsorption of oxygen may indeed be critical. Because the interest here is in oxidation of Sb_2O_3 in air, the role of oxygen was not examined.

To determine if a correlation between the equilibrium sublimation pressure of the antimony oxides and oxidative reaction rate should exist, if sublimation is rate determining, Langmuir's equation for sublimation was considered. (73)

In Langmuir's technique to determine the equilibrium sublimation pressure, $P_{\mathbf{e}}$, the weight loss during

evaporation is determined. The following relationship is then used to obtain the equilibrium sublimation pressure,

$$P_{e} = \frac{1}{m} \left(\frac{2\pi RT}{M} \right)^{1/2} \tag{4.3}$$

Where

P_e = equilibrium (e) sublimation pressure

m = rate of loss of mass per cm² sample per second

R = gas constant (1.987 cal/mole K)

T = sample temperature (K)

M = molecular weight of subliming species $(583.04g/\text{mole for } Sb_AO_6)$

q = accommodation coefficient

It is assumed in Langmuir's treatment that the accommodation coefficient, α , is equal to α_c , the condensation coefficient or fraction of molecules condensing per collision, and equal to α_s or the sublimation coefficient which is the ratio of the observed ratio of sublimation to the maximum possible rate of sublimation pressure. For reasonable values of α , values of m, close to experimentally determined values could be calculated.

Based upon Equation 4.3, the slope relating equilibrium sublimation pressure and mass loss is obtained as follows:

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$$\alpha \left(\frac{P_e}{m}\right) = \left(\frac{2\pi RT}{M}\right)^{1/2} \tag{4.4}$$

Thus, the slope deviates from linearity by a product of reaction temperature divided by the molecular weight of the subliming species to the half power which on inspection will be small over the oxidation temperatures investigated here. Therefore, if one simply plots m, the reaction rate for antimony trioxide oxidation, as a function of sublimation pressure, there should be an approximate linear correlation.

To determine if such a relationship exists for the oxidation of ${\rm Sb}_2{\rm O}_3$, reaction rate data gathered in the isothermal studies was analyzed.

Reaction rates (mg $0_2/g$ Sb $_20_3/sec$) were determined graphically from Figures 5 and 8. Sublimation pressures at those reaction temperatures were calculated using the appropriate relationships developed by Jungermann and Plieth. (50) Those data are presented in Tables 3 and 4 as follows:

Table 3. REACTION RATES AND SUBLIMATION PRESSURES FOR Sb203(0)

REACTION TEMP., K (°C)	REACTION RATE (mg 02/g Sb203/sec)	SUBLIMATION PRESSURE (torr)
729 (456)	0.00584	0.0188
766 (493)	0.01756	0.0788
779 (506)	0.02449	0.1295
795 (522)	0.04729	0.2244

Table 4. REACTION RATE AND SUBLIMATION PRESSURES FOR Sb203(c)

REACTION TEMP. K (°C)	REACTION RATE (mg O ₂ /g Sb ₂ O ₃ /sec)	SUBLIMATION PRESSURE (torr) Sb ₂ 0 ₃ (c)	SUBLIMATION PRESSURE (torr) Sb ₂ O ₃ (o)
794 (521)	0.01547	0.4894	
816 (543)	0.04784	1.0340	
832 (559)	0.06683	1.7378	0.7756
852 (579)	0.08449	3.2356	1.4411

Values of sublimation pressure for $\mathrm{Sb_2O_3}(o)$ at 832 and 852 K were also calculated because the reaction temperature of 832 K is just 3 K above the $\mathrm{Sb_2O_3}(c)$ to $\mathrm{Sb_2O_3}(o)$ transition temperature while 852 K is about 33 K above that transition.

Figure 16 is a plot of the $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ sublimation pressure versus the reactor rate. As seen in Figure 17, there is an equivalent plot for $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ oxidation. The sublimation pressure data plotted are for $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ at the lower two temperatures and for $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ at the highest temperature. For the reaction at 832 K, a mean (1.2567 torr) of the sublimation pressures for $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ and $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ is plotted. Again, it is evident that there is a linear dependence of the reaction rate upon the sublimation pressure.

Using least squares regression methodology, estimates of the appropriate regression coefficients were determined. For the reaction rate of ${\rm Sb_2O_3}(o)$ over the temperature range investigated, the least squares equation is as follows:

$$(\text{mg } 0^2/\text{g } \text{Sb}_2 0_3 (\text{o})/\text{sec}) = 0.1995 P_{\text{eq}} + 0.0013$$
 (4.5)
(correlation coefficient = 0.995)

For $Sb_2O_3(c)$, the equation is as follows:

$$(\text{mg } 0_2/\text{g } \text{Sb}_2 0_3(\text{c})/\text{sec}) = 0.0715 \text{ P}_{eq} - 0.0217$$
 (4.6)
(correlation coefficient = 0.995)

where

Peg = appropriate sublimation pressure in torr

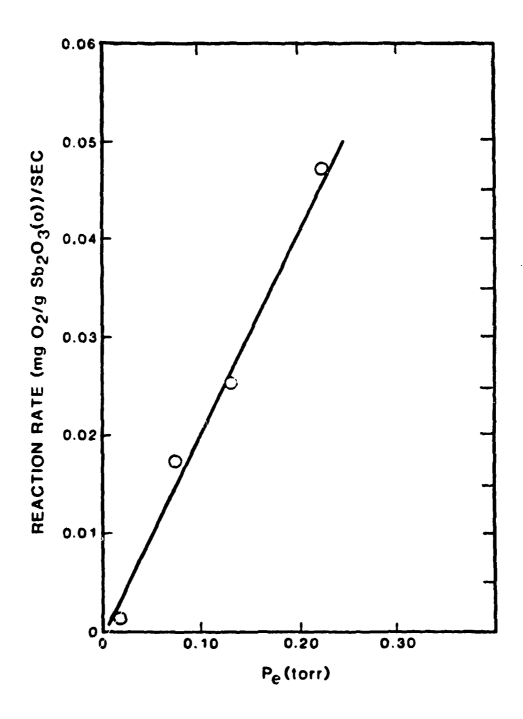


Figure 16. Oxidation Rate of ${\rm Sb}_2{\rm O}_3({\rm o})$ as a Function of Equilibrium Sublimation Pressure.

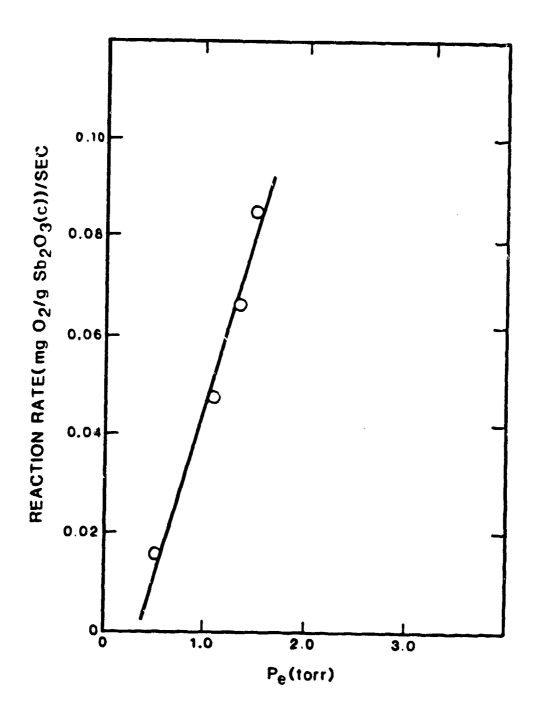


Figure 17. Oxidation Rate of Sb₂O₃(c) as a Function of Equilibrium Sublimation Pressure.

When the rate of oxidation of equimixed Sb_2O_3 , which is intermediate between pure $Sb_2O_3(o)$ and $Sb_2O_3(c)$ as presented in Table 5, is plotted as a function of mean sublimation pressure calculated using the Jungermann and Plieth⁽⁵⁰⁾ relationships, the correlation is not linear as data generated for the $Sb_2O_3(c)$ and $Sb_2O_3(o)$ reactions as seen in Figure 18. Comparing the oxidation rate at $520^{\circ}C$ (793 K) with that at $530^{\circ}C$ (803 K), there still appears to be a significant correlation of reaction rate with sublimation pressure, which supports the hypothesis that sublimation of Sb_4O_6 is the rate controlling step in oxidation of either $Sb_2O_3(o)$ or $Sb_2O_3(c)$.

Additionally, to determine if the free energy of oxidation for ${\rm Sb}_2{\rm O}_3$ is favorable for reaction to involve sublimed species, the following analysis was performed.

For the oxidation of antimony trioxide, two reactions must be considered. Those are:

$$Sb_2O_3(s) + 1/2 O_2 + Sb_2O_4(s)$$
 (4.7)

$$5b_40_6(g) + 0_2 + 5b_40_8(s)$$
 (4.8)

where ${\rm Sb_40_6}$ is the structure of the trioxide in the vapor state $^{(74)}$ and ${\rm Sb_20_4}$ is more correctly noted as ${\rm Sb_20_3}$. ${\rm Sb_20_5}$ or ${\rm Sb_40_8}$. $^{(28)}$

MIXED ${\rm Sb}_2{\rm O}_3$ OXIDATION RATES AND SUBLIMATION PRESSURE Table 5.

(torr)	HEAN	0.124	0.194	0.346	0.484
SUBLIMATION PRESSURE (torr)	sb ₂ 0 ₃ (o)	0.077	0.120	0.219	0.299
SUBLIMAT	sb ₂ 0 ₃ (c)	0, 171	0.267	0.473	0.668
	REACTION RATE [(mg02/gSb203)/sec]	5.42 x 10 ⁻³	6.02×10^{-3}	15.7 = 10-3	15.0×10^{-3}
	REACTION TEMP., K(°C)	765 (492)	777 (504)	793 (520)	803 (530)
	REACT	16!	11.	193	803

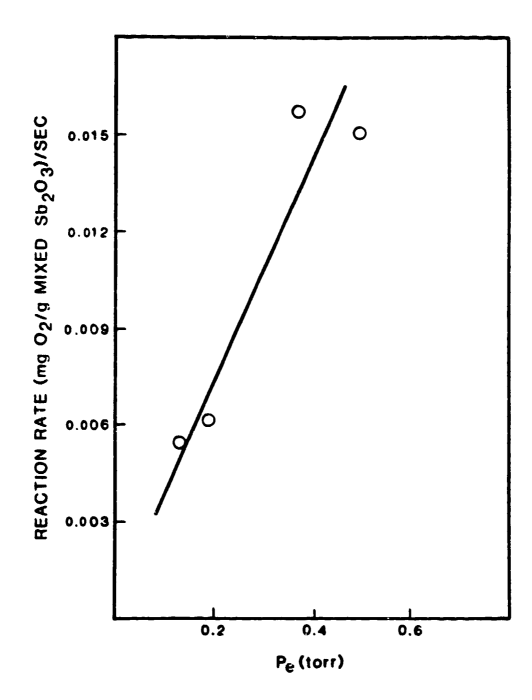


Figure 18. Oxidation Rate of Mixed Dimorphic Sb₂O₃ as a Function of Equi¹⁴ brium Sublimation Pressure.

By application of the free energy relationship,

$$\Delta G^{\circ} = -RT \ln K_{eq}$$
 (4.9)

where ΔG° = standard Gibbs free energy change for reaction at T

R = Gas constant = 1.987 cal/deg-mole

T = Temperature (K)

Keq = Equilibrium constant

It is possible to compare free energies of the reactions as follows:

For Equation 4.7,

$$K_{eq}(1) = \frac{[sb_2O_4]}{[sb_2O_3][p_{O_2}]^{1/2}}$$
(4.10)

and for Equation 4.8,

$$K_{eq}(2) = \frac{[Sb_4o_8]}{[P_{Sb_4o_6}][P_{O_2}]}$$
(4.11)

Standard states for $p_{0_2} = p_{Sb_40_6} = 1$ atm. For purposes of these calculations, the activities of $5b_20_3(s)$ and $5b_40_8(s)$ are assumed to be 1.

Values of ΔG^{\bullet} as a function of p_{0} for a reaction at 726°K with $Sb_20_3(o)$ sublimation pressure of 0.01648 torr

calculated using the Jungermann and Plieth relationship (50) are presented in Table 6.

Table . FREE ENERGIES FOR OXIDATION REACTIONS

	Equation 4.7	Equation 4.8
1.0	0	-15,490
0.5	- 500	-16,490
0.25	-1000	-17,490
0.184 (140 torr)	-1220	-17,930
0.1	-1660	-18,810
0.01	-3320	-22,140
0.001	-4980	-25,460

The values of ΔG° indicate that there is a much stronger driving force for the oxidation of the sublimed species. This is consistent with a mechanism involving sublimation of Sb_40_6 followed by reaction with 0_2 to form Sb_40_8 .

To determine the effect of sublimation pressure of Sb406 on ΔG° for Equation 4.8, calculations of ΔG° as a

function of temperature for P_{O_2} of 0.184 atm (140 torr) were made. Those data are presented in Table 7.

Table 7. FREE ENERGIES OF OXIDATION OF SUBLIMED Sb406

T(K) PS	b ₄ 0 ₆ equil (torr)	PSb ₄ O ₆ (atm) ln K _{eq} \(\Delta G^*(cal/mole) \)
726	0.01648	$21.68 \times 10^{-6} 12.432$ -17,930
766	0.07984	$105 \times 10^{-6} 10.854$ -16,520
779	0.1288	$169.5 \times 10^{-6} 10.375 -16,060$
794	0.2194	288.7×10^{-6} 9.843 -15,530

These data indicate that there is slightly less driving force for the reaction as the temperature is increased.

To compare the reaction tendency of $\mathrm{Sb}_2\mathrm{O}_3(c)$ with that of $\mathrm{Sb}_2\mathrm{O}_3(o)$, the sublimation pressure of $\mathrm{Sb}_2\mathrm{O}_3(c)$ is computed using the relationship developed by Jungermann and Plieth⁽⁵⁰⁾ and substituted into Equation 4.9. The results are given in Table 8.

Table 8. FREE ENERGIES FOR OXIDATION OF $Sb_2O_3(c)$ and $Sb_2O_3(o)$

Dimorph	T(E)	(etm)	(terr)	760	in the	AG*(cal/mole)
\$6 ⁵ 0 ³ (c)	298	0.184	4.29 m 10 ⁻²¹	5.65 x 10 ⁻²⁴	55.22	-32,700
	726	0.184	0.03639	4.79 x 10 ⁻³	11.64	-16,790
	794	0.184	0.40935	6.44 x 10 ⁻⁴	9.04	-14,260
\$6203(0)	298	0.184	2.26 x 10 ⁻²¹	2.97 x 10 ⁻²⁴	55.87	-33,080
	726	0.184	0.01648	2.17 x 10 ⁻⁵	12.43	-17,930
	794	9-184	0.2194	2.89 x 10 ⁻⁴	7.84	-15,530

The data indicate slightly less driving force for the reaction of ${\rm Sb}_2{\rm O}_3(c)$ in the 726-794 K interval compared with ${\rm Sb}_2{\rm O}_3(o)$ because of the higher sublimation pressure of the cubic dimorph at identical temperatures.

Based upon these analyses, the correlation of oxidation reaction rate with equilibrium sublimation pressure, and the previous observation that the reaction apparent activation energy is equivalent to the enthalpy of sublimation of the oxides, it appears that sublimation of Sb $_4$ 0 $_6$ is the rate controlling step for the oxidation of both Sb $_2$ 0 $_3$ (o) and Sb $_2$ 0 $_3$ (c). For Sb $_2$ 0 $_3$ (o), the proximity of a low energy reaction surface evidently increases the actual rate of reaction.

DIFFERENTIAL THERMAL ANALYSIS (DTA)

DTA was employed in additional thermal studies of antimony trioxides and MoS2. The DTA instrument is a

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Tracor-Stone Model 202 equipped with a Flatinel thermocouple system to measure sample and reference temperatures with an accuracy of ± 0.4 percent. NBS-ITCA Standard References Materials (Set 758) were routinely employed to confirm calibration. Typical 3-5 mg samples were heated at 10°C per minute under air or argon. Sample and references pans were nickel. Al₂0₃ was the reference material for all determinations. The Stone Model FIDF DTA furnace was also used in the isothermal oxidation studies.

Several DTA of Sb₂O₃ and other lubricant compact materials were completed. Those data are summerized in Appendix C. For samples of Sb₂O₃(o) from one lot ground for various lengths of times, it was found that as grinding time and hence surface area increased, an endothermic peak at about 650°-656°C associated with melting of Sb203 decreased in size. For the same samples, an exothermic peak around 485°-515°C associated with oxidation of Sb204 tended to increase in size, implying more complete oxidation, and decrease in peak temperature as grinding time increased. Figure 19 represents DTA 102-107 for 15 seconds, one and two minutes of grinding. Similar results were observed when Sb₂O₃(c) was ground, i.e., the temperature of peak oxidation decreased as grinding time increased. These data may be rationalized as oxidation of smaller particle size $\mathrm{Sb}_2\mathrm{O}_3(o)$ followed by melting and,

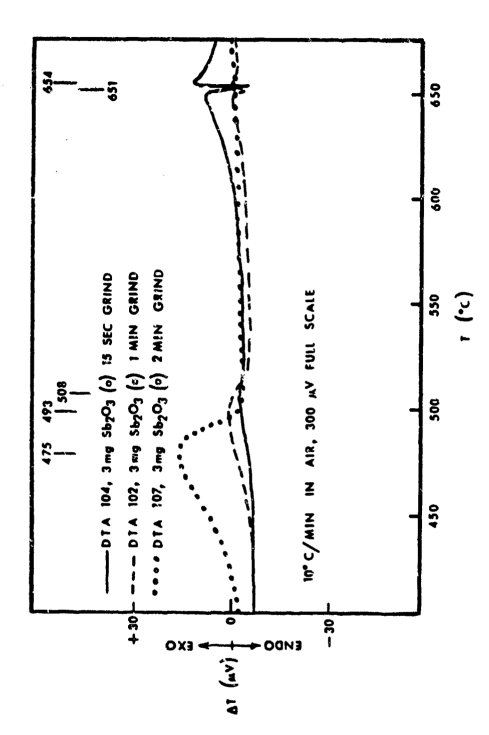


Figure 19. Effect of Grinding on Oxidation of $5b_20_3(\mathrm{o})$.

ultimately, oxidation of very large crystals of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$. Such grinding was, however, not reproducible from one lot of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ to another. For example, DTA 74 and 73 data from a different lot (#558649) of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ ground for eight and 16 minutes, respectively, were quite different from those for another lot as found in DTA 72 and 75. These results confirm the difficulty in reproducing particle size distribution in powders by simple grinding.

Because the effects of grinding ${\rm Sb}_2{\rm O}_3({\rm o})$ occurred as would be predicted, additional studies of the oxidation of pure dimorphic ${\rm Sb}_2{\rm O}_3$ by DTA was terminated.

For $\mathrm{Sb}_2\mathrm{O}_3(o)$ thermally analyzed under argon, no exothermic peaks were observed in the oxidation region as shown in Figure 20. A slight endothermic trend which was attributed to sublimation of $\mathrm{Sb}_2\mathrm{O}_3(o)$ was found through the typical oxidation region.

To determine the effect of $\mathrm{Sb}_2\mathrm{O}_4$ on the oxidation of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$, a mixture was prepared and analyzed by DTA. In Figure 21, the DTA are compared, where it is seen that addition of $\mathrm{Sb}_2\mathrm{O}_4$ actually shifts the oxidation peak to higher temperatures. Thus, the addition of $\mathrm{Sb}_2\mathrm{O}_4$ to $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ appears to act as a diluent for the exothermic oxidation of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$. For $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$, an addition of $\mathrm{Sb}_2\mathrm{O}_4$ did shift the oxidation peak to lower temperatures. To

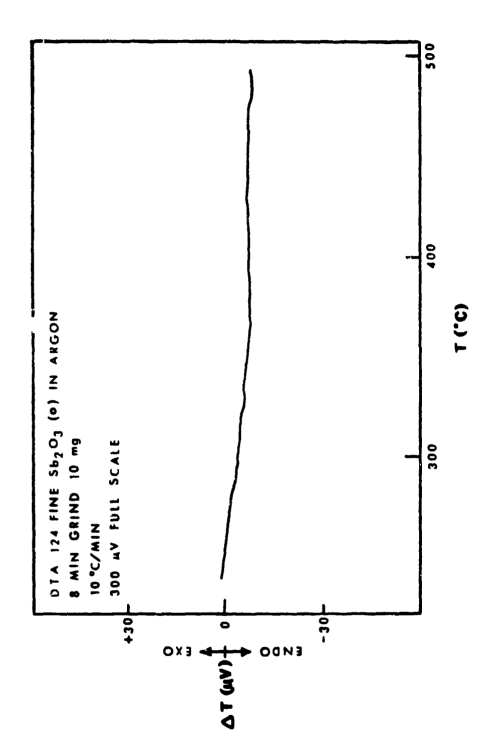
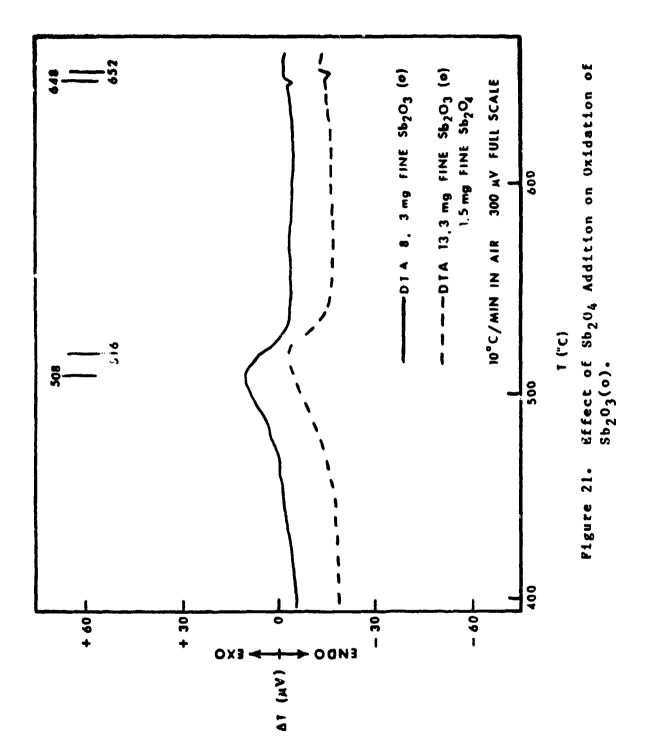


Figure 20. UTA of Sb203(o) Under Argon.

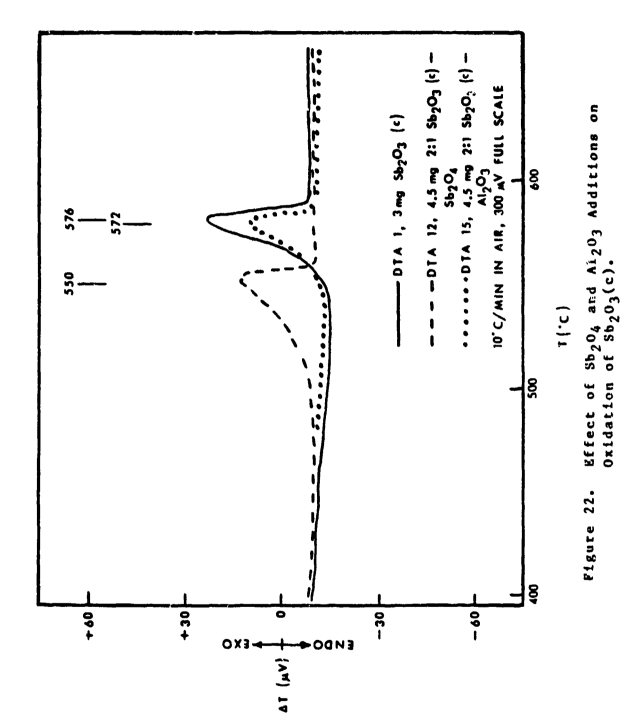


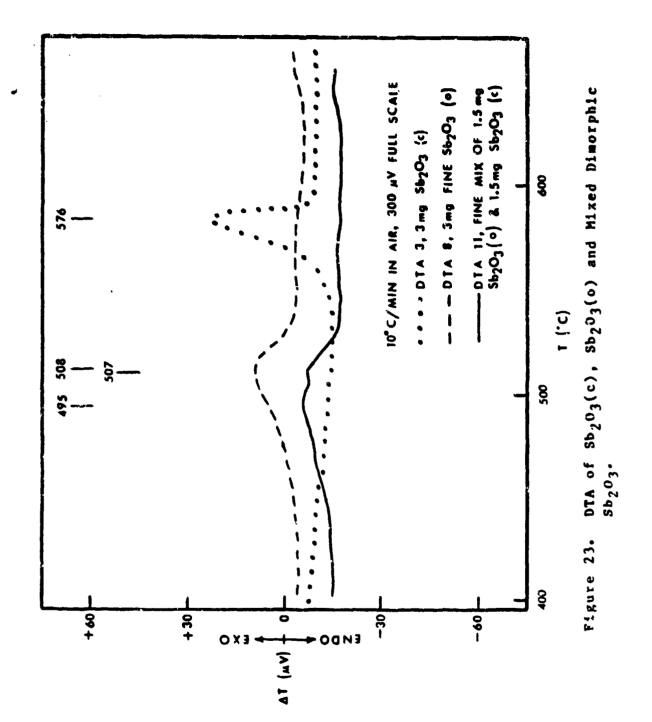
compare Sb_2O_4 with another diluent, Al_2O_3 was added to $Sb_2O_3(c)$ which did not significantly shift the oxidation peak. The area of the oxidation peak was, however, reduced, as shown in Figure 22.

In a similar fashion, DTA of a mixture of $Sb_2O_3(o)$ and $Sb_2O_3(c)$ was obtained and compared with DTA for $Sb_2O_3(o)$ and $Sb_2O_3(c)$. A composite of the DTA is presented in Figure 23. It is seen that oxidation of the mixture occurs at slightly below that of $Sb_2O_3(o)$ and significantly lower than $Sb_2O_3(c)$. The DTA for the mixed sample has two peaks which appears to be quite similar to very finely ground $Sb_2O_3(o)$ and may imply that some structure changes occur during the mechanical grinding treatment.

The results for mixed $\mathrm{Sb}_2\mathrm{O}_3$ suggest that $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ or $\mathrm{Sb}_2\mathrm{O}_4$ surface does influence the oxidation of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ as $\mathrm{Trofimov}$, et al. (72) suggest. However, $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ oxidation does not appear to be influenced by the presence of $\mathrm{Sb}_2\mathrm{O}_4$; in fact, $\mathrm{Sb}_2\mathrm{O}_4$ acts only as a diluent, which may result from the fact that $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ is orthorhombic itself. The size of the unground $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ is orthorhombic itself. The size of the unground $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ employed in mixed isothermal oxidation studies may be the influential factor in differences observed here and in those studies on the effect of mixing $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ and $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ on the rate of oxidation.

Additional DTA studies of mixed $\mathrm{Sb}_2\mathrm{O}_3$ samples were not pursued once it was established that there does appear to





be an effect of an orthorhombic surface on the oxidation of ${\rm Sh}_2{\rm U}_3(c)$.

To determine if the presence of $\mathrm{Sb}_2\mathrm{O}_3$ alters the oxidation of MoS_2 , DTA of MoS_2 , MoS_2 - $\mathrm{Sb}_2\mathrm{O}_3$ and $\mathrm{Sb}_2\mathrm{O}_3$ compacts were obtained. The DTA of powdered MoS_2 is presented in Figure 24. The onset of oxidation for MoS_2 is in agreement with data of Harmer and Pantano⁽⁸⁾ and $\mathrm{Peace}^{(75)}$. DTA of compacts of MoS_2 - $\mathrm{Sb}_2\mathrm{O}_3$ and $\mathrm{Sb}_2\mathrm{O}_3$ are given in Figure 25. As seen, the presence of $\mathrm{Sb}_2\mathrm{O}_3$ does not significantly influence the temperature of MoS_2 oxidation. These data suggest that there is not a role for antimony oxide in the oxidative protection of MoS_2 . Concurrent wear studies presented later provided evidence that there is no significant anti-oxidant role for $\mathrm{Sb}_2\mathrm{O}_3$ in MoS_2 compacts, and this phase of the study was concluded.

SUMMARY OF Sb203 OXIDATION STUDIES

Isothermal oxidation studies of the oxidation under 140 torr exygen sublimation is rate controlling for oxidation of either $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ or $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$. The oxidation of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ appears to require higher reaction temperatures for rates equivalent to $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$. Mixtures of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ and $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ appeared to be intermediate between $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ and $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$; however, larger sized $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ was used inadvertently in the mixtures than in $\mathrm{Sb}_2\mathrm{O}_3$ exidations. Since reaction rates involving sublimation are highly dependent upon surface area, the isothermal data for mixed

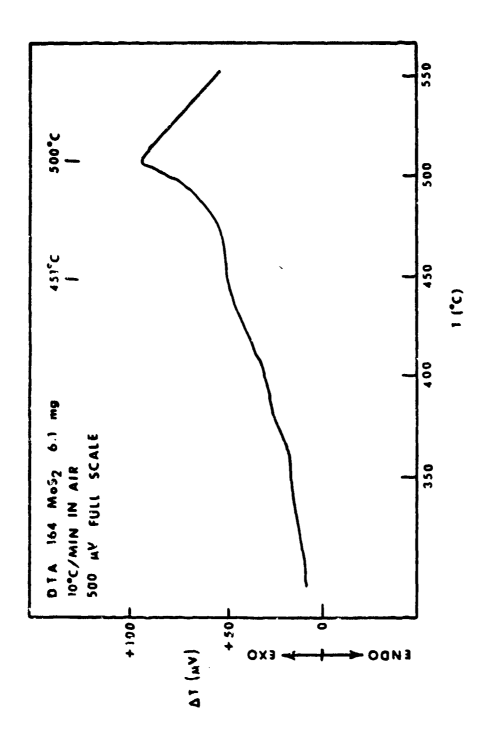
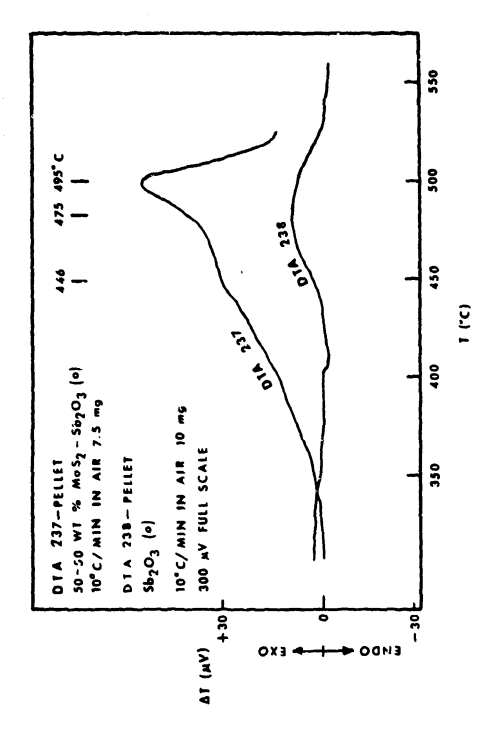


Figure 24. DTA of MoS2.



Pigure 25. DTA of Compacts of MoSz-Sb203 and Sb203.

 ${\rm Sb}_2{\rm O}_3$ supports a mechanism of oxidation which includes sublimation but does not confirm the observations of Trofimov, et al. $^{(72)}$ The role of sublimation in oxidation of ${\rm Sb}_2{\rm O}_3$ is supported by calculations of free anargies of oxidation for ${\rm Sb}_2{\rm O}_3$ (s) and the sublimed species: ${\rm Sb}_4{\rm O}_6$, which indicate a greater driving force for the oxidation of ${\rm Sb}_4{\rm O}_6$.

DTA results, however, indicate that mixtures of ${\rm Sb}_2{\rm O}_3({\rm c})$ with ${\rm Sb}_2{\rm O}_3({\rm o})$ or ${\rm Sb}_2{\rm O}_4$ do oxidize at lower temperatures supporting the observation of Trofimov and coworkers. DTA oxidation of ${\rm Sb}_2{\rm O}_3({\rm o})$ or ${\rm Sb}_2{\rm O}_3({\rm c})$ is in agreement with data of Golunski, et al. (56) Additionally, DTA of ${\rm MoS}_2$ confirms that oxidation begins at about 300°C and is completed near 500°C. When a compact of ${\rm MoS}_2{\rm -Sb}_2{\rm O}_3$ is analyzed, there is no significant difference in peak temperature for oxidation of ${\rm MoS}_2$.

These data do not support a role for $\mathrm{Sb}_2\mathrm{O}_3$ in retarding oxidation of MoS_2 nor as sacrificially oxidizing instead of MoS_2 since $\mathrm{Sb}_2\mathrm{O}_3$ does not initially oxidize at temperatures comparable with the onset of oxidation of MoS_2 .

CHAPTER V

PROPERTIES OF MoS2-Sb203 SOLID LUBRICANT COMPACTS

Several properties of $MoS_2-Sb_2O_3$ compacts are critical to development of an understanding of that lubricating system. Areas of prime interest are how the compacts form, how they wear and what changes occur at the interface. The significance of each must be established to provide a basis for understanding of the beneficial mechanism of Sb_2O_3 additions to MoS_2 .

COMPACTION

To establish if compaction of Sb₂O₃-MoS₂ plays an atypical role in its superior performance, previous efforts in the area were reviewed. In that review, it was found that Harmer and Pantano⁽⁸⁾ compacted cylinders of MoS₂ with 25 and 45 wt% of Sb₂O₃ in a laboratory press to examine densification of solid lubricant compacts. Those data have been reduced and summarized in Table 9. The data were then employed to calculate the theoretical density fractions attained by compaction in Table 10. Those theoretical compact densities were plotted as a function of forming pressure in Figure 26. From Figure 26, several observations are apparent:

DEMSITY OF COMPACTS FOR DIFFERENT FORMULATIONS AND FORMING PRESSURES* Table 9.

COMPOSITION

Wt. 2	Vol. 2	(12)	FORMING PRESSURE (MPa)	RE (MPa)
		PELLET		DENSITY (g/cm3)**
		172	345	069
100% MoS ₂	190	3.147	3.453 3.455	3.740
75% MoS ₂ 25% Sb ₂ 0 ₃	77.26	3.319	3.636 3.605	3.940
55% MoS ₂ 45% Sb ₂ 0 ₃	58.05 41.95	3.796	3.751 3.709	4.032
A Compute Assumin	ed from Reference 8 data. ng densities of 4.80 g/cm 57 g/cm ³ for Sb ₂ 0 ₃ (o) and	Computed from Reference 8 data. Assuming densities of 4.80 g/cm 3 for MoS $_2$, 5.2 g/cm 3 for Sb $_2$ 0 $_3$ (c) and that Sb $_2$ 0 $_3$ used was a equimolar	MoS2, 5.2 g/cr Sb ₂ 0 ₃ used waa	n^3 for $Sb_2O_3(c)$ s a equimolar

mixture of the Sb203 dimorphs.

THEORETICAL COMPACT DENSITY PRACTION FOR DIFFERENT FORMULATIONS AND FORMING PRESSURES* Table 10.

0.7792 0,7946 0.7928 069 THEORETICAL DENSITY FRACTION PORMING PRESSURE (MP&) 0.7196 0.7333 0.7301 0.6556 0.6693 0.746 172 Vol. X 58.05 41.95 77.26 22.74 100.0 COMPOSITION 75% MoS₂ 25% Sb₂0₃ 55% MoS₂ 45% Sb₂0₃ 100% MoS2

* Computed from Reference 8 data.

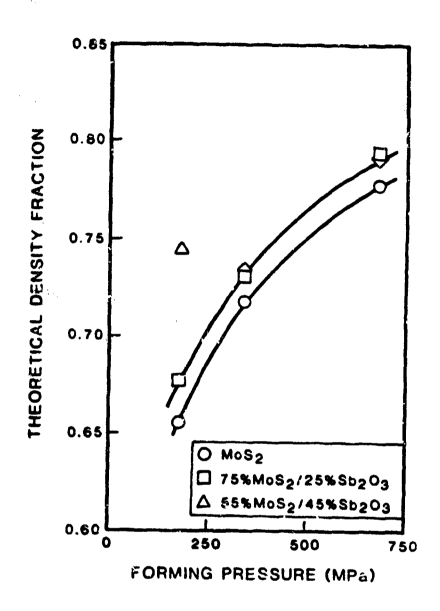


Figure 26. Theoretical Density Fraction as a Function of Forming Pressure from Harmer and Pantano. (8)

- (i) Under the highest load for all compacts, only about 80% of theoretical density was obtained.
- (ii) Both ${
 m MoS}_2$ and ${
 m MoS}_2$ containing 25 wt% ${
 m Sb}_2{
 m O}_3$ followed a similar pattern in that there was a gentle increase in fraction of theoretical density achieved as a function of increasing forming pressure.
- (iii) For MoS₂ containing 45 wt% of Sb₂O₃, about 75% of theoretical density was achieved at the lowest pressure, or approximately 8% more than the compacts containing less Sb₂O₃. It also should be noted that the data for the intermediate pressure does not follow the expected trend. It is not known if there was a formulation, pressure, or other error or if the data presented are within a normal error distribution. Thus, it would be concluded that the addition of Sb₂O₃ aids in compaction and/or densification of solid compacts based upon their results. It is also seen that it might be possible to attain unusually high densification at lower pressures under optimum conditions if the low pressure, high Sb₂O₃ concentration data were valid.

To validate the previous compaction data of Harmer and Pantano $^{(8)}$, high purity $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ and MoO_3 (Alfa, Danvers, MA) were individually ground mechanically and sieved through a 30 $\mu\mathrm{m}$ nickel screen. Oxides were weighed and thoroughly mixed with weighed technical grade MoS_2 (Bemol, Elmwood, CN) which is more adequately described in a later

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section or wear of MoS₂-Sb₂O₃. These pellets were then compacted in a cylindrical die mounted in a laboratory press. Forming pressures were about 172 MPa (25,000 psi).

On removal from the die, compacts were measured and densities computed. Those data for each compact are presented in Table 11. The ratio of apparent density to theoretical density is a function of compact composition is given in Figure 27. It was found in these experiments that the addition of Sb₂O₃ or MoO₃ to MoS₂ does not improve compaction at relatively modest pressures at ambient. trend for the MoS2-Sb2O2 compacts is the opposite of that found by Harmer and Pantano. The opposite trends found are difficult to rationalize, although differences in oxide particle size may be significant. However, the largest difference is for pure MoS2 where at similar forming pressure, Harmer and Pantano only achieved about 83% of the apparent density for MoS2 obtained in this work. Although the difference in trends was obvious, efforts were not pursued to resolve the conflict since the experiments in this work indicate that large additions of $\mathrm{Sb}_2\mathrm{O}_3$ do not promote a typical densification of MoS2 compacts at low pressures. Rather efforts were directed towards determining the influence of $\mathrm{Sb}_2\mathrm{O}_3$ and MoO_3 on compaction of MoS2 at high temperatures.

COMPACTION OF MoS AND MOS WITH Sb203 OR MOO3 AT AMBIENT Table II.

FFLLET	PELLET	PELLET DIAMETER (cm)	PELLET HEIGHT (cm)	DENSITY (g/cm ³)	P _E Theoretical Density (g/cm³)	ما م	MEAN 9 9 T
NoS ₂	0.5017	0.041	0.413	3.786	30 30	0.776	
	0.4758	0.641	0.388	3.800		6.776	6.771 ±3.001
22.13 vol X NoO ₃ 77.87 vol X NoS ₂	0.4865	0.641	0.431	3.498	4	0.723	
41.1 vol Z Noû3 58.9 vol Z Nos ₂	0.4823	0.641	0.457	3.265	08.4	0.680	
23.05 vol % 8b ₂ 0 ₃ 0.4823 76.95 vol % Hos ₂	0.4823	0.641	0.394	3.792	5.01	0.757	
42.33 vol X Sb203 0.4939	0.4939	0.641	0.404	3.785	5.12	0.739	
57.67 vol Z Sb ₂ 0 ₃ 0.7628	0.7628	0.641	0.623	3.813		0.745	0.74
	0.7707	0.641	0.618	3.864		0.755	*0.00

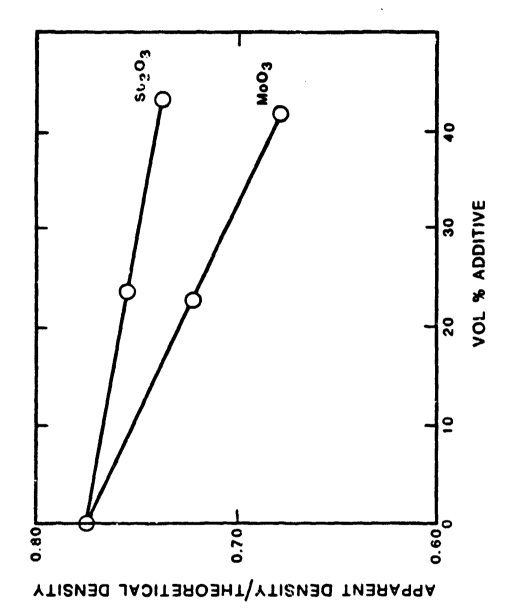


Figure 27. Density Ratio as Function of Volume Percent Additive.

To determine thermal expansion and softening properties of MoS₂ and MoS₂ plus additives at temperature, compressed cylinders were prepared in a 0.635 cm (0.25 in.) diameter die and pressed at 172.4 MPs. (25 ksi). The TMA employed a counterweighted loading system so it was possible to load all samples at one gram. The flat load rod was 2.48 mm in diameter and 4.83 mm² in contact area resulting in an extremely low pressure of 2.93 kPa (0.32 psi). The TMA increase in temperature was 2°C/win for all samples which were inerted under nitrogen.

For an $\mathrm{Sb}_2\mathrm{O}_3$ cylinder evaluated from room temperature to about 650°C under nitrogen, there appeared to be several regions of interest as the cylinder expanded in response to the increasing temperature. First, there is a region of modest expansion (15.5 $\mu\text{m/m/°C}$) from ambient to about 225°C after which the slope approached zero up to about 350°C. The rate of expansion then increased (43.1 $\mu\text{m/m/°C}$) in the approximate temperature range of $400^\circ-510^\circ\text{C}$. Again, there was a region of reduced slope followed by a period of very rapid expansion near the $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})-\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ transition. The expansion peaked at 610°C followed by rapid compression. These chans in the rate of thermal expansion for the polycrystalline compact are attributed to expansion of the crystal lattice, conversion of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ to $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ followed by fusion of $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$.

For a MoS₂ cylinder, reduction in cylinder height was found from ambient to about 375°C. For the 7.57 mm length cylinder, the reduction in dimension was about 45 μ m over that temperature interval. From 375°C to 769°C, the cylinder increased in length at a modestly accelerating rate (mean of ~ 40 μ m/m/°C). Above 769°C, the cylinder again compressed until the end of the test at about 780°C.

The early compression of the ${\rm MoS}_2$ cylinder is attributed to reorganization of the crystallites to a more stable structure. The pronounced effect in ${\rm MoS}_2$ is thought to originate from the layered structure which would contribute to a strongly unisotropic structure initially formed in the die.

For a 50-50 weight percent sample of $\mathrm{Sb}_2\mathrm{O}_3$ and MoS_2 , the expansion was virtually nil until about 300°C. To about 353°C, the structure expanded at a rate of about 45.8 $\mu\mathrm{m/m/°C}$. Then expansion was very limited until 440°C, when rapid expansion took place which peaked at an estimated temperature region of 560°-570°C. The TMA results for the MoS_2 - $\mathrm{Sb}_2\mathrm{O}_3$ compact are given in Figure 28.

The expansion results for the ${\rm Sb}_2 {\rm O}_3$ - ${\rm MoS}_2$ cylinder are thought to be the result of balancing the expansion of ${\rm Sb}_2 {\rm O}_3$ with the contraction of ${\rm MoS}_2$ up to about 350°C followed by mutual expansion up to the power of fusion of ${\rm Sb}_2 {\rm O}_3$.

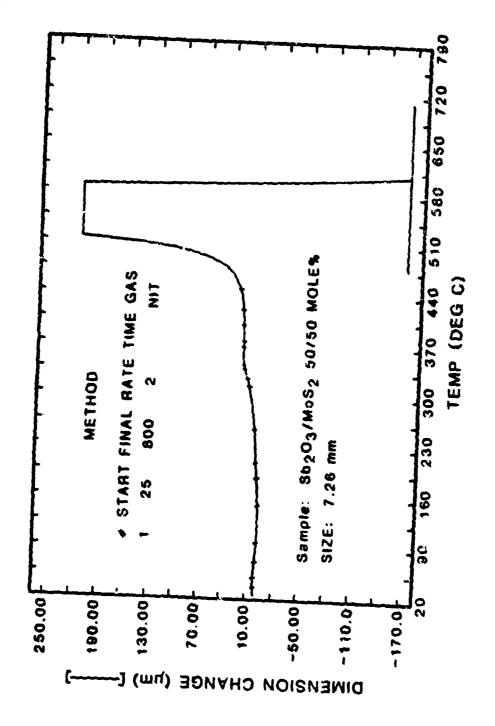


Figure 28. THA of Mos₂-sb₂0₃ Compact.

A cylinder of 25 mole percent Sb203-25 mole percent MoO3-50 mole percent MoS2 was evaluated by TMA. The sample was initially heated under nitrogen to about 160°C when the TMA inadvertently held at that temperature until restarted. It was found that at about 275°C, the cylinder began to expand rapidly. After another 100°C temperature rise, the cylinder had expanded more than 5 µm longer than the original length of 8.49 mm, after which it was not instrumentally possible to record the rapid expansion. about 725°C, the length of the cylinder had been sufficiently reduced so that the TMA was able to record the reduction in length. The expansion was attributed primarily to the presence of MoO₃ in the Sb₂O₃-MoS₂ compact. The Sb203-Mc03-MoS2 pellat began expanding significantly at a temperature about 175°C lower than the Sb203-MoS2 pellet.

WEAR OF MoS2-Sb203 COMPACTS

To study the effects of additions of $\mathrm{Sb}_2\mathrm{O}_3$ on the wear life of compacts, wear experiments were conducted at several temperatures under varying conditions. Compacts were formed and shaped for testing using the materials and procedures described below. Molybdenum sulfide was of technical grade from Bemol, Elmwood, CN; with composition and particle size specifications as given in Table 12.

Table 12. DESCRIPTION OF MOS2 FOR COMPACTS

COMPOSITION	WEIGHT PERCENT	PARTICLE SIZE	WEIGHT PERCENT
MoS ₂	98.2	On US Steve No. 100 (>150 µH)	0
Acid Insoluble	0.35	Through 100 on 200 (<150 to >75 µH)	2
Iron	0.15	Through 200 or 325 (<75 µM to >45 µM)	10
Molybdenum Trioxide	0.01	Through 322 (<45 µH)	85
Water	00.00		
011	0.03		
Carbon	1.00		
Acid Number	0.01		

The molybdenum disulfide is representative of what is available commercially for solid lubricant applications and was used without modification.

The antimony oxides were prepared identically to those described previously for oxidation studies. Because of the size of $Sb_2O_3(o)$ formed in the furnace, it was necessary to finely grind and sieve the product before compaction (<20 µm). Initially, it was thought that by simply sieving $Sh_{>}O_{3}(c)$ through a 325 mesh screen (<44 µm) would be sufficient, however, it was found later that it was necessary to grind Sb₂O₃(c) also to obtain the least wear. The effects of Sb₂O₃(c) particle size on wear shall be discussed in a subsequent section. Materials were weighed to the nearest tenth of a milligram with a total cylinder weight of 300 mg. The compacts were prepared by thoroughly mixing the weighed components and pressing at 275.2 MPa (40 ksi) in a cylindrical die in a laboratory press. The compact mounted in the die was trimmed by lathe. The angle of cut was 21.8 degrees receding from the face. The substrates were 440C stainless steel disks with a surface hardness of 57-60 Rc.

Wear disks were prepared by initial light, wet sanding with 400 grit followed by 600 grit silicon carbide paper.

Lapping was performed using a Lapmaster Model 12 machine employing 3 micrometer corundum powder. Disks were cleaned with Stoddard solvent and rinsed in water followed by

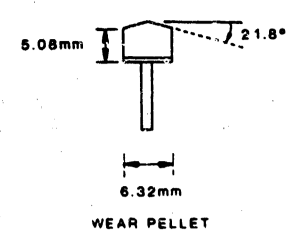
isopropyl alcohol prior to oven drying at approximately 60°F. A surface finish of 4-6 microinches was attained as measured by a Bendix Profilometer, Type OB, Model 18. The profilometer was calibrated using a glass roughness standard.

Tripellet evaluation of solid lubricant compositions were performed initially using the Midwest Research Institute (MRI) pellet wear-life machine. (76) The machine as pictured in Figure 29 with the wear cone and mounting schematic shown in Figure 30, permits loading three lubricant compacts with a total load of 300 g. Sliding velocity is approximately 3.81 m/s (750 ft/min).

The frictional force generated during cliding was measured by use of a linear variable differential transformer (LVDT) mounted to measure deflection of a spring loaded retainer. The LVDT output was recorded by strip chart recorder. Run times were typically 20 minutes at room temperature, but test times were modified as necessary to achieve measurable cone dismeters. Since cones were used, comparisons between tests must be based upon identical tests including time and sliding distance so that wear volume data may be useful. It is reasonable to assume that the wear equation such as the one proposed by Archard for adhesive wear and described previously, (14)

$$V = \frac{k L \pi}{3p} \tag{5.1}$$

Figure 29. Triplellet Wear Test Machine.



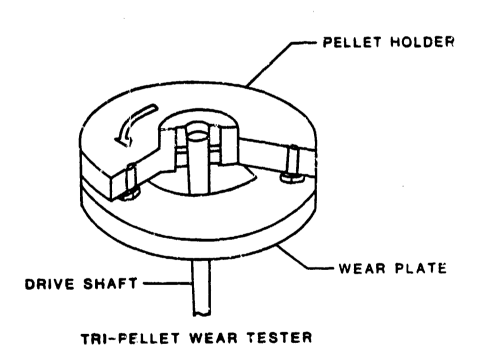


Figure 30. Wear Cone and Pellet Mount Configuration.

The wear volume, V may be calculated as follows:

$$V = \frac{\pi}{3} r^2 h = 1.047 r^2 h \tag{5.2}$$

and for a 21.8° cone,

$$h = 0.39997 r$$
 (5.3)

then substituting,

 $V = (1.047)(0.39997\pi)r^2$

 $= 0.4188 \text{ m}^3$

r = radius of worn cone base

h = worn cone height

After microscopic measurement of the cone diameter, wear volume was then calculated using the above relationship. The microscope was a Bausch and Lomb (25x). A stage micrometer was used for calibration.

Pellet loads were computed based upon the weight of the mount, holder and solid lubricant compact. Of course, as the pellet wears, the load is reduced insignificantly; however, the apparent contact stress decreases significantly during wear tests because the pellet contact area increases, so data may be used for comparative purposes only.

Some tripellet tests conducted at room temperature were performed with the test machine installed in a drybox manufactured by Vacuum/Atmosphere Co., Hawthorne, CA (Model HE-403 Dri-Train). Water removal was accomplished by use of a molecular sieve. The drybox atmosphere may be continuously circulated through the molecular sieve bed using a fan and duct system.

The concentration of atmospheric moisture was varied for wear of MoS₂-antimony oxide compacts. Trace moisture content was determined with a Shaw hygrometer manufactured in Bradford, England. The instrument permits the measurement of trace atmospheric moisture at the part per million level.

To obtain very high levels of moisture in the drybox, the atmosphere was saturated by bubbling air through water and passing through the drybox.

Only limited wear tests were conducted because a great amount of effort was required to prepare materials for each specimen and to evaluate the compacts.

TRIPELLET WEAR TESTS AT ROOM TEMPERATURE

Dry Air

Initial testing conducted in a drybox substantiated the dramatic effect of atmosphere upon MoS_2 wear properties, which has been reviewed previously. (77) In four hour tripellet tests on 440C steel substrates in very dry air (1-3 ppm H_2O), wear of MoS_2 was comparable with

MoS₂ containing 25 mole percent of Sb_2O_3 or Sb_2O_4 at about 0.2 to 0.3 mm³ per compact with minor effects of the oxide additions found. Very low friction coefficients (=0.05) were generated in most evaluations with the exception of $MoS_2-Sb_2O_4$ at 0.1-0.13. (Tests 2-9).

Wet Air

When the test environment was saturated with H2O, it was necessary to shorten test times to one minute for MoS₂. The wear volume (3.9 mm³) for MoS₂ in wet air was about an order of magnitude larger than tests that were 240 times longer in dry air (Tests 2 and 10). Data generated in the tests (Tests 11-16) indicates that NoS2 with Sb₂O₃(o) or Sb₂O₃(c) wore less than compacts of MoS₂. A compact of MoS2-Sb204 wore at a rate of about three times greater than compacts containing Sb2O3, but wore at a much slower rate than pure MoS2 compacts. Friction coefficients for all compacts were uniformly high (0.1-0.33) and highly irregular in saturated air. Inspection of compacts by optical and scanning electron microscopy suggest that the oxides were providing a stable support structure in a soft matrix of MoS2. Discrete antimony oxide particles could be found around which MoS, appeared to be flowing.

SINGLE PELLET WEAR TESTS

For the next series of wear tests, the single pellet MRI Mark-VB machine (76) was employed as shown in Figure

The tests (Tests 29-35) were conducted in air introduced through desiccant into the test chamber (~50-80 ppm H₂O, except Test 35 because of a moisture leak) at room temperature for 20 minutes. With the Mark-VB apparatus, it was not possible to reduce H₂O levels to the level achieved with earlier tests in a dry box. The data are summarized in Table 13. The wear volumes were smaller than earlier tripellet tests conducted in either very dry or very wet atmospheres. The lowest wear exhibited was with MoS2- $Sb_2O_3(o)$. In fact, $Sb_2O_3(o)$ was the only beneficial additive. Sb203(o) was also tribologically superior in earlier drybox tests but not to the extent seen in these tests. The larger wear volumes found for Sb₂O₃(c) were later attributed to the bipyrimidal crystal structure of $Sb_2O_3(c)$, which, even though sieved through 325 mesh (< 44 µm), appeared to abrade wear surfaces. Lowest coefficients of friction were the lowest for MoS2-Sb203(o) formulations. MoS2-Sb204 compacts initially generated coefficients comparable with $MoS_2-Sb_2O_3(o)$ but increased substantially during the test. Mechanical problems precluded the measurement of friction coefficients for MoS2~Sb2O3(c) compacts. These data indicated a superiority for compacts containing \$b203(0).

WEAR EVALUATIONS AT 160°C

Five tests were run at 160°C for 10 minutes: two (Tests 37 and 38) in embient air (50% RH), and three (Tests

Table 12. SINGLE PELLET WEAR VOLUMES

RT, Dry Air, 20 Minutes, 440C Wear Disks

TEST NUMBER	PELLET COMPOSITION (Mole Percentage)	WEAR VOLUME (mm ³)	COEFFICIENT OF FRICTION
30	MoS ₂	0.17	0.1 - 0.13
31	75 MoS_2 - 25 Sb_2O_3 (o) 75 MoS_2 - 25 Sb_2O_3 (o)	0.05	0.06 - 0.07
35	75 $MoS_2 - 25 Sb_2O_3(c)$. 75 $MoS_2 - 25 Sb_2O_3(c)$	2.46	
33	75 MoS ₂ - 25 Sb ₂ 0 ₃ 75 MoS ₂ - 25 Sb ₂ 0 ₄	0.24	0.06 - 0.13 0.05 - 0.17

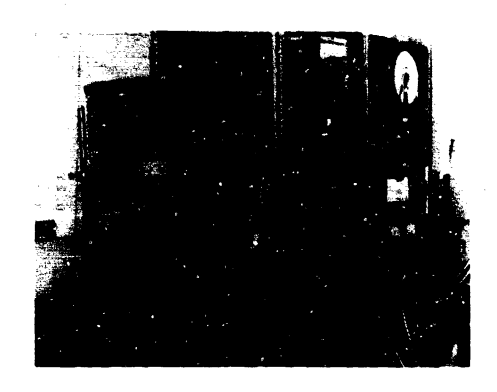


Figure 31. MRI Mark-VB Single Pellet Wear Test Machine.

39-41) in dry argon (<80 ppm 0_2). Those data are presented in Table 14. It was found that MoS_2 compacts with 25 mole percent $\text{Sb}_20_3(o)$ generated less wear volume than pure MoS_2 compacts under both conditions. These data suggest that the role of Sb_20_3 in improvement of MoS_2 tribological properties does not involve oxygen or oxidation. Interpretation of friction data is difficult in that the values determined are closely grouped.

WEAR EVALUATIONS AT 232°C

An important observation is that the wear volume of compacts containing $\mathrm{Sb}_2\mathrm{O}_3(c)$ that was ground and sieved identically to $\mathrm{Sb}_2\mathrm{O}_3(o)$ produced much lower wear volumes than compacts containing unground $\mathrm{Sb}_2\mathrm{U}_3(c)$. Comparing wear data and scanning electron microscopy of pellets with both ground and unground $\mathrm{Sb}_2\mathrm{O}_3(c)$ suggests that sieved unground $\mathrm{Sb}_2\mathrm{O}_3(c)$ can abrade the wear surface much more easily than ground $\mathrm{Sb}_2\mathrm{O}_3(c)$, because the unground particles tend to form masses of abrasive materials compared with ground

Table 14. SINGLE PELLET WEAR DATA AT 160°C

CUEPPICIENT OF PRICTION AIR ARGON 502 RH	0.09 - 0.11 0.09 - 0.12 0.09 - 0.10	0.08 - 0.10
WEAR VOLUME (mm ³) AIR ARGON 502 RH	0.13	0.09
COMPOSITION (Mole Percent)	MOS ₂ Mos ₂ Mos ₂	MoS ₂ - 25 Sb ₂ O ₃ (o) 0.09 MoS ₂ - 25 Sb ₂ O ₃ (o)
₹ aT NUMBER	37 39 41	38 75 1

WEAR EVALUATIONS AT 232°C ON 440C HEAR DISKS Table 15.

Hos2 Hos3 Hos3		COMPOSITION	WEAR	WEAR VOLUME (mm ³)		FRICTION CORFFICIENT	CIENT
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		(Hale Percent)		WET AIR AR	2	VET AIR	ARGON
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	9	Mo S ₂		0	35		0.11 - 0.12
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	47	MoS	91.0		0.12		
75 MoS ₂ - 25 Sb ₂ 0 ₃ (o) 0.02 0.06 - 0.07 0.06 - 0.07 0.08 0.06 0.09 0.03 0.03 0.08 - 0.09 0.03 0.03 0.03 0.09 0.03 0.03 0.03	52	Ho S ₂		0.20		0.13 - 0.14	
75 $hos_2 = 25 sb_2o_3(o) 0.02$ 75 $hos_2 = 25 sb_2o_3(o) 0.13$ 75 $hos_2 = 25 sb_2o_3(o) 0.13$ 76 $hos_2 = 25 sb_2o_3(c)$ 77 $hos_2 = 25 sb_2o_3(c)$ 78 $hos_2 = 25 sb_2o_3(c)$ 79 $hos_2 = 25 sb_2o_3(c)$ 71 $hos_2 = 25 sb_2o_3(c)$ 72 $hos_2 = 25 sb_2o_3(c)$ 73 $hos_2 = 25 sb_2o_3(c)$ 74 $hos_2 = 25 sb_2o_3(c)$ 75 $hos_2 = 25 sb_2o_3(c)$ 75 $hos_2 = 25 sb_2o_4$ 76 $hos_2 = 25 sb_2o_4$ 77 $hos_2 = 25 sb_2o_4$ 78 $hos_2 = 25 sb_2o_4$ 79 $hos_2 = 25 sb_2o_4$ 71 $hos_2 = 25 sb_2o_4$ 71 $hos_2 = 25 sb_2o_4$ 72 $hos_2 = 25 sb_2o_4$ 73 $hos_2 = 25 sb_2o_4$ 74 $hos_2 = 25 sb_2o_4$ 75 $hos_2 = 25 sb_2o_4$ 76 $hos_2 = 25 sb_2o_4$ 77 $hos_2 = 25 sb_2o_4$ 78 $hos_2 = 25 sb_2o_4$ 79 $hos_2 = 25 sb_2o_4$ 70 $hos_2 = 25 sb_2o_4$ 70 $hos_2 = 25 sb_2o_4$ 71 $hos_2 = 25 sb_2o_4$ 72 $hos_2 = 25 sb_2o_4$ 73 $hos_2 = 25 sb_2o_4$ 74 $hos_2 = 25 sb_2o_4$ 75 $hos_2 = 25 sb_2o_4$ 76 $hos_2 = 25 sb_2o_4$ 77 $hos_2 = 25 sb_2o_4$ 78 $hos_2 = 25 sb_2o_4$ 79 $hos_2 = 25 sb_2o_4$ 70 $hos_2 = 25 sb_2o_4$ 71 $hos_2 = 25 sb_2o_4$ 72 $hos_2 = 25 sb_2o_4$ 73 $hos_2 = 25 sb_2o_4$ 74 $hos_2 = 25 sb_2o_4$ 75 $hos_2 = 25 sb_2o_4$ 76 $hos_2 = 25 sb_2o_4$ 77 $hos_2 = 25 sb_2o_4$ 80 $hos_2 = 25 sb_2o_4$	42	NoS ₂ - 25 8b ₂	•	•0	95		0.08
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75 MoS ₂ - 25 Sb ₂ 0 ₃ (c) 0.13 0.07 - 0.09 75 MoS ₂ - 25 Sb ₂ 0 ₃ (c) 2.2 75 MoS ₂ - 25 Sb ₂ 0 ₃ (c) 1.78 0.07 - 0.10 75 MoS ₂ - 25 Sb ₂ 0 ₃ (c) 0.39 (Ground) 0.06 - 0.07 75 MoS ₂ - 25 Sb ₂ 0 ₄ 75 MoS ₂ - 25 Sb ₂ 0 ₄ 76 MoS ₂ - 25 Sb ₂ 0 ₄ 77 MoS ₂ - 25 Sb ₂ 0 ₄ 78 MoS ₂ - 25 Sb ₂ 0 ₄ 78 MoS ₂ - 25 Sb ₂ 0 ₄ 79 MoS ₂ - 25 Sb ₂ 0 ₄	6 77	HoS ₂ - 25 Sb ₂				0.08 - 0.09	
75 Mos ₂ - 25 Sb ₂ 0 ₃ (c) 2.2 2.1 0.03 - 75 Mos ₂ - 25 Sb ₂ 0 ₃ (c) 1.78 0.07 - 0.10 75 Mos ₂ - 25 Sb ₂ 0 ₃ (c) 0.39 (Ground) 0.06 - 0.07 75 Mos ₂ - 25 Sb ₂ 0 ₃ (c) 0.39 (Ground) 0.06 - 0.07 75 Mos ₂ - 25 Sb ₂ 0 ₄ 6.31 0.23 0.08 - 0.11 75 Mos ₂ - 25 Sb ₂ 0 ₄ 0.31 0.02 0.08 - 0.11 75 Mos ₂ - 25 Sb ₂ 0 ₄ 0.31 0.02 0.03 0.07 - 0.13 75 Mos ₂ - 25 Sb ₂ 0 ₆ 0.31 0.27 0.09 - 0.09 - 0.09 - 0.00 0.00 0.00 0.0	57	5 MoS ₂ - 25 Sb ₂				•	
75 HoS_2 - 25 $\text{Sb}_2\text{O}_3(c)$ 1.78 0.07 - 0.10 75 HoS_2 - 25 $\text{Sb}_2\text{O}_3(c)$ 0.39 (Ground) 0.06 - 0.07 75 HoS_2 - 25 $\text{Sb}_2\text{O}_3(c)$ 0.39 (Ground) 0.06 - 0.07 75 HoS_2 - 25 Sb_2O_4 0.31 0.08 - 0.11 0.08 - 0.11 75 HoS_2 - 25 Sb_2O_4 0.17 (Ground) 0.09 - 0.09 - 0.09 -	43	Hos ₂ - 25 Sb ₂	(2.	2		0.07 - 0.12
75 MoS ₂ - 25 Sb ₂ O ₃ (c) 1.78 0.39 (Ground) 0.06 - 0.10 1.78 0.35 - 25 Sb ₂ O ₃ (c) 0.39 (Ground) 0.06 - 0.07 1.75 MoS ₂ - 25 Sb ₂ O ₄ 0.31 0.23 0.08 - 0.11 0.07 - 0.13 1.75 MoS ₂ - 25 Sb ₂ O ₄ 0.31 0.27 0.13 0.07 - 0.13 1.75 MoS ₂ - 25 Sb ₂ O ₄ 0.17 (Ground)	4.5	MoS ₂ - 25 Sb ₂	~	2.	-		
75 $\text{MoS}_2 - 25 \text{ Sb}_2 0_3 \text{(c)}$ 0.39 (Ground) 0.06 - 0.07 75 $\text{MoS}_2 - 25 \text{ Sb}_2 0_4$ 0.31 0.23 0.07 - 0.13 75 $\text{MoS}_2 - 25 \text{ Sb}_2 0_4$ 0.17 (Ground) 0.07 - 0.13 75 $\text{MoS}_2 - 25 \text{ Sb}_2 0_4$ 0.17 (Ground)	55	Hos ₂ - 25 Sb ₂	~	1.78		0.07 - 0.10	
75 MoS ₂ - 25 Sb ₂ 0 ₄ 6.31 0.08 - 0.11 75 MoS ₂ - 25 Sb ₂ 0 ₄ 6.31 0.27 0.27 0.07 - 0.13 75 MoS ₂ - 25 Sb ₂ 0 ₄ 0.17 (Ground)	56	5 MoS ₂ - 25 86 ₂			(pun	0.06 - 0.07	
75 $MoS_2 - 25 Sb_2O_4$;	NoS ₂ - 25		•0	23		0.07 - 0.10
75 MoS ₂ - 25 Sb ₂ 0 $_4$ 0.27 0.13 75 MoS ₂ - 25 Sb ₂ 0 $_4$ 0.17 (Ground)	64	Hos2 - 25 Sb2	6.31		0.08 - 0.1	•	
75 MoS ₂ - 25 Sh ₂ O ₄ 0.17 (Ground)	54	Mos ₂ - 25 Sb ₂		0.27		0.07 - 0.13	
	69	Mos2 - 25 Sb2		0.17 (Gro	(pun		0.09 - 0.11

 ${\rm Sb}_2{\rm O}_3(c)$. These data invalidate earlier test data generated here at lower test temperature from pellets containing unground ${\rm Sb}_2{\rm O}_3(c)$. Grinding ${\rm Sb}_2{\rm O}_4$ in a similar manner reduces compact wear volume. The grinding permits more intimate mixing with ${\rm MoS}_2$ and reduces the probability that large abrasive particles will damage the substrate solid lubricant film.

Additionally, Lavik, Hubbell and McConnell⁽¹⁾ indicate that MoS₂ with additions of Sb₂O₃ are superior tribologically in both air and vacuum to MoS₂ compacts, strongly inferring that a role for oxidation of MoS₂ or Sb₂O₃ is non-existent. Limited wear studies of compacts presented in Table 15 in dry air and dry argon confirm that Sb₂O₃ addition to MoS₂ is beneficial for wear in both air and inert atmosphere at 232°C. Thus, a beneficial role involving oxidation for additions of Sb₂O₃ to MoS₂ cannot be supported.

SINGLE PELLET WEAR EVALUATIONS AT 316°C

All tests at 316°C were performed on the MRI Mark-VB machine. 440C steel wear disks were used for 10 minute tests 61 to 73; all other evaluations employed Type 302 stainless steel shim stock (0.05 mm thick) with test times of 1000 seconds.

The first series of evaluations at 316°C was to generate data for comparison with results from the tests at lower temperatures. This series of tests is summarized in

Table 16. At 316°C, the effects of atmosphere are apparent. Under argon, the wear is typically greater than for identical evaluations in air. These results confirm the effects of the atmosphere as reported in the literature, (77) where it is noted that MoS₂ wear in argon is high because MoO₃ does not form. MoO₃ is believed to bind MoS₂ crystals together for improved performance.

These results at 316°C indicate that antimony oxides improve the tribological performance of MoS_2 . Results with $Sb_2O_3(c)$ are believed to be the result of large, unground crystals of $Sb_2O_3(c)$ which effectively abrade the interface. The significance of the shape and size of $Sb_2O_3(c)$ was only established after these initial experiments at 316°C.

MoS₂ compacts containing $Sb_2O_3(o)$ yielded not only lower wear volumes compared with pure MoS_2 , but also lower coefficients of friction. $Sb_2O_3(o)$ wear was marginally reduced in argon compared with air. Because previous tests showed that $Sb_2O_3(o)$ additions were most beneficial, wear tests with $Sb_2O_3(c)$ and Sb_2O_4 were not repeated.

Similar tests were performed on the stainless steel shim stock previously described, primarily to generate wear surfaces that could be easily mounted for examination by scanning and transmission electron microscopy. Those data are summarized in Table 17. The data indicated that 25

Table 16. INITIAL WEAR STUDIES At 316°C

TEST #		A FOR A OLD OR CHES	E (mm,)	FRICTION C	PRICTION CORPFICIENT
62	(Mole Percent)	DRY AIR	ARGON	DRY AIR	ARGON
6.5	Mo S ₂	0.36	• •	0.14 - 0.15	
7.0	Mo S ₂	91.0	• • •	0.08 - 0.14	0.10 - 0.11
61 75 66 75	$1008_2 - 25 8b_20_3(0)$ $1008_2 - 25 8b_20_3(0)$	0.19	0.12	0.07 - 0.09	0.07 - 0.09
63 75 68 75	$MoS_2 - 25 Sb_2O_3(o)$ $MoS_2 - 25 Sb_2O_3(o)$	7.97	5.6	0.08 - 0.2	0.05 - 0.14
64 75	MoS ₂ - 25 Sb ₂ 04 MoS ₂ - 25 Sb ₂ 04	0.18	0.54	0.08 ~ 0.19	0.08 - 0.19

Table 17. INITIAL ${\sf MoS}_2$ antihony oxide wear tests on 0.05 mm SS shim stock in dry air (100 g load) at 316°C

	COMPOSITION		
TEST	(Nole Percent)	WEAR VOLUME (mm3)	COEPFICIENT OF PRICTION (µ)
7.6	MoS ₂	0.35	
79 75 78	65 MoS ₂ - 35 Sb ₂ 0 ₃ (o) 75 MoS ₂ - 25 Sb ₂ 0 ₃ (o) 85 MoS ₂ - 15 Sb ₂ 0 ₃ (o)	0.20 0.13 0.32	0.07
76	75 MoS ₂ - 25 Sb ₂ O ₃ (c)	1.12	
11	75 HoS ₂ - 25 Sb ₂ 04	0.25	0.21 - 0.36
8	75 HoS ₂ - 12.5 Sb ₂ 0 ₃ (o) - 12.5 Sb ₂ 0 ₃ (c)	0.12	0.07 - 0.10

mole percent of ${\rm Sb_2O_3(o)}$ or a total of 25 mole percent of equinized ${\rm Sb_2O_3(o)-Sb_2O_3(c)}$ are effective additives for ${\rm MoS_2}$ under these conditions.

A simple evaluation of Sb₂O₃(o) concentration (Tests 75, 78, 79) is also given in Table 17 suggesting that the 25 mole percent concentration is better than 15 or 35 mole percent. However, as with many wear tests, the data are only single points, and substantial conclusions cannot be made; rather, trends were established to identify areas for additional effort.

EFFECT OF LOAD ON WEAR IN 316°C TESTS

Tests were also conducted to determine if variation of wear volume as a function of load for $MoS_2 - Sb_2 O_3(o)$ compacts required any additional study. The results as given in Table 18 generated expected results, i.e., wear was reasonably proportional to load, and the effort was terminated.

EXAMINATION OF WEAR SURFACES BY ELECTRON MICROSCOPY

An attempt was made to examine wear tracks (Wear Tests 73-76) produced on the 0.05 mm stainless steel substrates using transmission electron microscopy (TEM). Using the standard preparative technique of embedding the sample in an organic resin, curing and cutting thin sections with a diamond edge, it was found that samples suitable for TEM could not be prepared. Evidently, the very soft wear track

Table 18. EFFECT OF LUAD ON WEAR VOLUME FOR TESTS AT 316°C FOR 1000 SECONDS

FRICTION	60,		•
COEFFICIENT OF	.0709	8 8	90.
LOAD WEAR VOLUME (mm ³) COEFFICIENT OF FRICTION	0.05	0.13	0.17
LOAD (R)	80	100	150
COMPOSITION (Mole Percent)	75 HoS ₂ - 25 Sb ₂ 0 ₃ (o)	75 HoS ₂ - 25 Sb ₂ 0 ₃ (o)	75 MoS ₂ - 25 Sh ₂ 03(0)
TEST	82	75	83

materials varied so greatly in hardness from the steel foil that uniform cross-sections could not be removed as a whole.

When the foils were mounted in a standard metallurgical mount and polished, for scanning electron microscopic examination, the results were disappointing. Again, disturbance of the wear track and blending of that section into the mounting material appeared to limit the usefulness of the technique.

Wear compacts and tracks were also prepared for direct examination by scanning electron microscopy (SEM) in the following manner. Tracks and compacts were coated with a layer of silver to improve conductivity of the sample. Coating was done in a Ernest F. Fullum Vacuum Evaporator Model 12500, at a vacuum of 10⁻³ torr. The specimens were placed 10 cm from the tungsten basket which contained silver wire that was evaporated under vacuum at a current of less than 20 amperes.

The samples were examined in an Amray SEM Model 1600. Samples were positioned perpendicular to the electron beam at an accelerating voltage of 30 keV and a 2 distance of 12 mm for accurate measurement capability. Since all conditions remained the same, the relative sizes of worn area could be estimated from photographs made at the same magnification.

at 316°C have a rough appearance compared with MoS₂ compacts containing Sb₂O₃ evaluated under the same conditions. For the MoS₂ sample, after wear at 316°C for 1000 seconds in dry air, the compact surface contained cracks and fissures as well as an abundance of wear debris (Wear Test 115). A comparative test of a wear compact (Test 116) containing Sb₂O₃ under the identical test conditions produced a much smoother surface. This is evident in SEM views of the two surfaces in Figures 32 and 33.

This observation is in agreement with that of Harmer and Pantano $^{(8)}$ who found that densification and sintering of worn compacts of $MoS_2-Sb_2O_3$ was obvious compared with MoS_2 wear compacts.

EUTECTIC FORMATION OF Sb2O3 WITH MOO3

To determine if a sutectic of $\mathrm{Sb}_2\mathrm{O}_3\mathrm{-MoO}_3$ is formed during sliding of $\mathrm{MoS}_2\mathrm{-Sb}_2\mathrm{O}_3$ compacts, and to elucidate a possible beneficial role of such a eutectic in those compacts, a eutectic mixture was prepared and compared with materials formed on sliding of $\mathrm{MoS}_2\mathrm{-Sb}_2\mathrm{O}_3$ compacts.

In the hypothesis advanced by Lavik, Hubbell and Meconnell, (7) eutectics formed from bulk additives and oxides produced during sliding, e.g., Sb_2O_3 with MoO $_3$ from MoS $_2$, would improve the tribological process. The

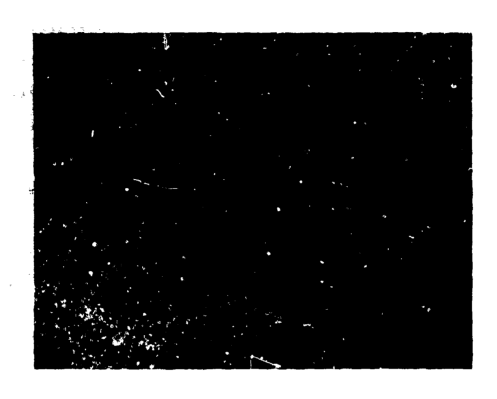


Figure 32. SEM Micrograph of MoS2 Worn Surface Compact.

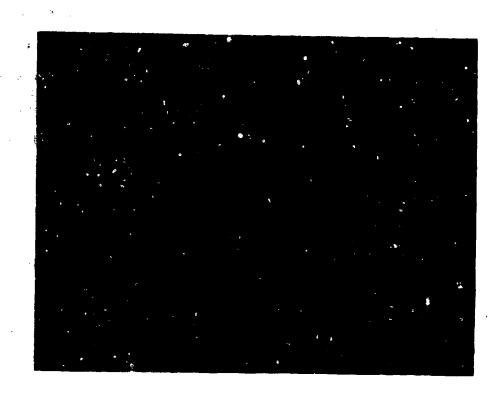


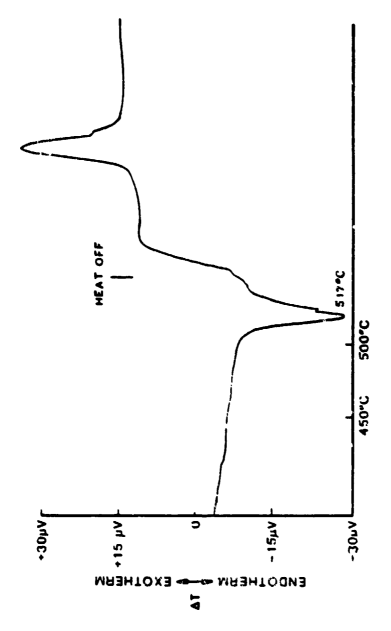
Figure 33. SEM Micrograph of MoS₂-Sb₂O₃ Worn Compact Surface.

mechanism of improvement was not investigated; however, those authors speculated that oxides bond to substrate surfaces which might improve attachment of the lubricant, i.e., act ss "bonding" agents for MoS2. The virtue of eutectic formation was spaculated to be that oxide weldment and attachment would be possible or could occur at lower temperatures. At the time of that work, data had not been published on the eutectic formation between Sb203 and MoO₃. However, in 1975, Parmentier, et al. (64) did publish the results of an investigation of \$b203-MoO3 eutectic formation. Those authors found that when mixtures of antimony oxide and molybdenum oxide were mixed and sealed under argon in glass ampules and heat-treated at 500°C for 24 hours, then eutectics were formed. While Sb2O3 melts at $656\,^{\circ}\text{C}$ and MoO_3 melts at 795 $^{\circ}\text{C}$, three eutectic temperature minima were detected. The temperatures of the eutectic points are 518°, 536° and 596°C. The minima were determined by use of differential thermal analyses of the heat-treated materials. Diffraction patterns of Sb2MoO6 and $Sb_2(MoO_4)_3$ compositions were obtained using x-ray techniques.

Parmentier, et al. data indicate that indeed eutectic formation between ${\rm Sb}_2{\rm O}_3$ and ${\rm MoO}_3$ is possible and that such materials in concentrations as required to produce the lowest melting combination is conceivable on the sliding interfaces of ${\rm MoS}_2{\rm -Sb}_2{\rm O}_3$ compacts.

To investigate the possibility for a role of the \$b₂0₃-MoO₃ eutectic in the improvement of the tribological performance of MoS2-Sb2O3 compacts, finely ground compositions of 25 mole percent Sb₂O₃-75 mole percent MoO₅, and 75 mole % Sb₂O₃-25 mole % MoO₃ were mixed and added to quartz tubes. The tubes were then evacuated (10^{-6} torr) three times and filled with 99.998% argon. On the fourth evacuation, the tubes were refilled to a pressure of 150 torr. The tubes were then separately inserted into a muffle furnace and allowed to stand for 24 hours at 500 ± 10°C. During the early phases of heat-treating, the tube containing the highest concentration of MoO2 shattered. This experiment was not repeated. For the other composition, the tube was opened after air quenching. A sample of the material was examined using differential thermal analysis. 75 mole percent Sb₂O₃-25 mole percent MoO₂ mixture, in a differential thermogram was obtained using the Stone DTA. An endotherm was detected at 517°C for 10 mg sample heated at 10°C/min. under argon. When the DTA furnace was allowed to cool, an exotherm of about the same size and shape was observed as shown in Figure 34. Thus, the findings of Parmentier, et al. were confirmed in that a eutectic of Sb203-MoO3 was produced and is detectable.

MoS2-Sb203 compacts were then worn at ambient to generate surfaces suitable for x-ray diffraction and DTA



ure 34. DTA of MoO3-Sb203 Eutectic Composition

examination to determine if a eutectic is formed. Compacts of 25 mole percent $Sb_2O_3(o)$ and 75 mole percent HoS_2 were prepared in cylinders rather than the standard conical shape and worn in the tri-pellet test three hours at 850 rpm.

Ten milligram samples of the worn surface areas of the compacts worn in air at room temperature on the tripellet machine were mechanically removed from the pellets and examined by DTA. Under argon, a sharp endotherm indicating the presence of a eutectic was not observed; rather, a small gentle endotherm was observed. The endotherm found is similar to that for small particle size $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ analysed by DTA under argon. These data indicated that if a eutectic of $\mathrm{Sb}_2\mathrm{O}_3$ -MoO $_3$ was formed at the pellet interface, then the concentration is so small as to be non-detectable using the DTA method.

Those areas and unknown areas of compacts were also analyzed using the technique of symmetric reflection wide angle x-ray diffraction (WAXD). Two instruments were employed, i.e., a Philips automatic powder diffractometer and Picker FACS-1 four-circle single crystal diffractometer. The Philips system used nickel filtered Cu Ka radiation at 35 kV and 22 ma. The worn compacts surface was scanned at either 1°/min. or 1/8°/min. The continuous scintillation detector output was recorded by a strip chart recorder. The Picker data was collected

digitally at 0.1° increments from selected areas (1 mm x 1 mm) of a sample. The Picker employed Cu Ko radiation at 30 kV and 20 mÅ a with an incident beam quartz crystal monochromator.

Operation of the Philips unit followed standard factory procedure. The Picker operation (78) and analysis (79) are described elsewhere.

X-ray diffraction data for non-worn and worn areas were compared. For each diffraction angle, the difference in counts was calculated and a spectrum of the differences plotted as given in Figure 35. Experimental x-ray data for values of 20, where 0 is the angle of diffraction, were reduced using the Bragg relationship: (80)

 $n\lambda = 2d \sin\theta \tag{5.4}$

where

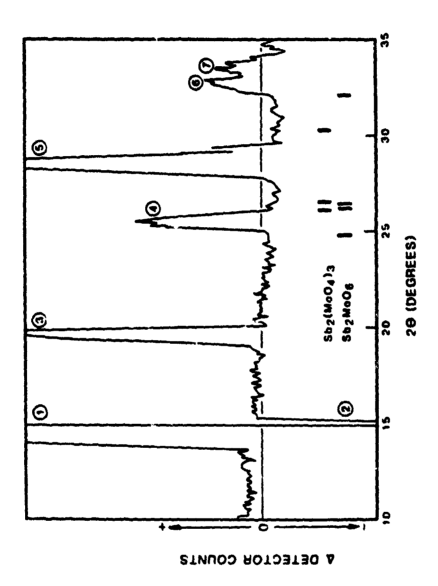
n = Order of diffraction

 λ = Incident radiation wavelength

d = Distance between crystal lattice atomic planes

 θ = Diffraction angle.

In the non-worn areas, the more prominent diffraction peaks listed in Table 19 were found to be stronger than in the worn area.



Offferential X-ray Offfraction Pattern for Worn and Non-Worn compact Surface. Figure 35.

Table 19. X-RAY DIFFRACTIONS MORE PROMINENT

IN THE NON-WORN AREAS

DIFFRACTION PEAK	d(Å)	ASSIGNMENT
IN FIGURE 35		
1	6.17	MoS ₂
3	4.52	Sb ₂ 0 ₃ (0)
4	3.49	sb ₂ 0 ₃ (o)
5	3.13	st ₂ 03(0)
6	2.72	\$b ₂ 0 ₃ (0)
7	2.67	\$b203(o)

The spectra contained diffraction patterns coincident with those of ${\rm MoS}_2$ and ${\rm Sb}_2{\rm O}_3(o)$. (81)

The only diffraction peak found in worn areas at about 5.90Å and is attributed to broadening of the \mbox{MoS}_2 difraction.

 ${
m MoS_2-Sb_2O_3}$ compacts were worn at 371°C using a small drill press to rotate on a temperature controlled hot surface. Again, DTA and x-ray analysis did not detect the presence of a cutectic of ${
m MoO_3-Sb_2O_3}$. However, a

ᡧᢔᡐᢓᡭᡪ᠘ᡟᡭᡛᢓᡭ᠘ᡥᡦᡛᠹᡭᠻᠼᠻ᠘ᡛᡚᢣᡦ᠘ᢊ᠘ᡛ᠘ᡀᢗᢖ᠘ᡀ᠘ᡀ᠘ᡀ᠘ᡀᢓᡀᢓᡀᢓᡀᡭᡀᡭᡀᡭᡀᡭᡀᡭᡀᡭᡀᢓᡀᢤ᠘ᡀ᠘ᡀ᠘ᡀ᠘ᡀ᠘ᡀ᠘ᡀ᠘᠘᠘᠘᠘᠘᠘᠘

significant change was the transformation of $\mathrm{Sb}_2\mathrm{O}_3(c)$ to $\mathrm{Sb}_2\mathrm{O}_3(o)$. This transformation which occurred under interface temperatures, provides direct evidence for contact temperatures of nearly 200°C in excess of test temperature since the transition temperature for $\mathrm{Sb}_2\mathrm{O}_3$ is 556°C. Evidence was also found for trace levels of $\mathrm{Sb}_2\mathrm{O}_4$ indicating some oxidation of $\mathrm{Sb}_2\mathrm{O}_3$ occurs under these conditions.

The lack of DTA and x-ray diffraction evidence indicates that eutectic compositions are not formed at the interface.

CHAPTER VI

DEVELOPMENT OF HYPOTHESIS FOR MoS2-Sb2O3 SOLID LUBRICANT COMPACTS

Studies of $\mathrm{Sb}_2\mathrm{O}_3$ have shown that the rates of oxidation of both $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ and $\mathrm{Sb}_2\mathrm{C}_3(\mathrm{c})$ to $\mathrm{Sb}_2\mathrm{O}_4$ are dependent upon surface area and the presence of an orthorhomic surface such as $\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ or $\mathrm{Sb}_2\mathrm{O}_4$. It was found that the presence of MoS_2 or MoO_3 did not materially affect the rate of $\mathrm{Sb}_2\mathrm{O}_3$ oxidation in compacts.

It was also found that although a eutectic of Sb₂O₃/MoO₃ could be produced in the laboratory as other workers had done, ⁽⁶⁴⁾ no evidence of eutectic formation on sliding interface; produced at ambient or at 371.2°C (700°F) was detected.

Wear studies of $MoS_2-Sb_2G_3$ compacts from ambient to 315.6°C (600°F) confirmed studies of earlier investigators, e.g., high relative humidity or the absence of oxygen in the environment increases wear of both MoS_2 and $MoS_2-Sb_2G_3$ compacts in room temperature evaluations. Under the same conditions, $MoS_2-Sb_2G_3$ formulations were superior to MoS_2 only if particle size of the antimony oxide additions was small. For Sb_2G_3 (c) additions, if the particle size were large, the relative

wear rate was high and was attributed to its larger bipyramid crystals abrasively ploughing and damaging the sliding interface. Under dry air conditions, with MoS2-Sb203 peliet wear specimens containing small partiale size SbyOg(o), the wear track and compact surfaces produced in tests were routinely dense and coherent in appearance compared with other materials evaluated, including pure MoS2 compacts. At higher temperatures, it was again found that wear compacts composed of Sb203 and MoS2 were superior to test specimens of MoS2 in that smoother frictional traces were produced, lower coefficients of friction were found, and wear volumes generated in identical tests were reduced. Also, MoS2-Sb503 formulations were superior to MoS2 whether evaluated in inert or oxygen-containing atmospheres. These experiments indicate that a role for Sb₂O₃ in MoS₂ in which Sb₂O₃ imparts superior tribological performance by acting in any way as an antioxidant or suppressing oxidation of MoS2 cannot be supported.

Based upon this analyzis, an alternative hypothesis is developed. The hypothesis is that Sh₂O₃ softens, i.e., plastically deforms under load and temperature conditions of this test so that it acts as a matrix to retain a preferred orientation of the primary solid lubricant crystallites, MoS₂, generated at asperity contact. Localized heating of the mixture permits softening of

the antimony oxide around and away from the sliding interface so that the lowest energy orientation of MoS_2 would be effectively "frozen in place" because asperity contact temperature rapidly recedes as sliding continues. This concept is supported by observations of others, (82) who state that "dense" wear tracks and/or test specimen surfaces are produced when the oxide is present. The concept is also supported by previous efforts (7) which found that the tribologically superior performance of MoS2-Sb2O2 compacts compared wih MoS2 specimens is independent of atmospheric oxygen content. Those authors alluded to an optimum formulation (7) which complements the hypothesis developed in that an optimum amount of addicive is required to permit efficient construction of the best tribological surfaces; too much additive results in the matrix material acting as the solid lubricant, diminishing the tribolobical properties of the mixture because of reduced primary solid lubricant content.

The concept suggests that the additive, Sb_2O_3 , should not be unique in its ability to improve MoS_2 performance. Also, bulk additions of stable materials properly selected should improve the performance of other solid lubricants. Again, data and observations of others support the hypothesis. For example, $DiSapio^{(83)}$ notes that the basis for improved tribological properties of MoS_2 in air versus in vacuum is that in air MoO_3 forms, which acts as a binder

for MoS₂. Similarly, in 1955, Peterson and Johnson (84) found that additions of GdO to graphite improved the tribological performance of graphite when tested up to temperatures of 537.8°C (1000°F).

A possible role of oxides in compacts can be seen in a study of the effectiveness of inorganic oxides as primary solid lubricants (85) in two standard four-ball machines. (86) The authors found that there was excellent correlation of the load-wear index as a function of the logarithm of Vickers hardness, equivalent to a correlation with Mohs hardness, (87) of those oxides. They concluded that the several oxides provided improved extreme pressure capability for a lubricant and suggested that the effect was a purely mechanical one, with better performance for the soft oxides. Softness of an oxide leads to tribological improvement because the oxide can then deform in a highly-loaded contact to prevent or reduce metal-tometal contact, thereby lowering the contact loads at ball surfaces. These results indicate that the ability of an oxide to deform under tribological conditions is a critical factor in their beneficial performance.

This study suggests that hardness, shear strength and melting point or decomposition temperature are critical criteria in selecting additives in this concept. It also indicates that test conditions are equally important criteria, since tribological and material properties are

strongly dependent on temperature, load, etc. If one considers a correlation of hardness and melting point for oxides, (88,89) it would appear that there are limited choices for bulk additions once test conditions are known. The additive of choice will be one that is stable under the test conditions and softens effectively at asperity flash temperature.

Examination of these data also indicate that only those materials with very low hardness and relatively very high melt point are natural solid lubricants over a wide range of conditions. Consider, for example, that molybdenum disulfide has a Mohs hardness of 1.5 with a melting point of 1185°C, and that graphite has a Mohs hardness of 0.5 to 1.0 with a sublimation temperature of approximately 3650°-3700°. (28) The softness of these materials results from a layer lattice structure. These materials and a few other structural analogs are the only known natural solid lubricants over wide temperature ranges. Other solid lubricants not possessing such structure which permits the atypical low hardness-high temperature stability are evidently only useful in a narrow temperature range under specified test conditions. In that limited range, the solid is soft or perhaps even liquid permitting shear with only modest forces involved.

Test conditions are important in the evaluation of the solid as a lubricant. For example, in a pin on disk

experiment, flash temperatures may typically be 200°C over the experimental temperature. Often, those temperatures become extremely high, e.g., from 1000° to 1200°C for steel and ceramic pins slid over sapphire (90). Under those conditions it is easily visualized that the material deposited on a wear track may resolidify, and if the hardness and/or structure permit the residue to become abrasive, then the solid will be lowly ranked. If, however, the experiment is designed so that cooling cannot occur on the disk, e.g., the ends of two equal diameter shafts operating in counter-rotation, then the lubricant would be ranked higher because it would lubricate effectively at the higher overall temperature.

Peterson, et al., (91) found that there is a correlation between coefficients of friction and melting points at a given test temperature, better for some classes of materials than others. For example, the friction coefficient of Na₂WO₄ at 704°C is 0.17. Friction coefficients of other tungstates and mclybdates increased as the difference between melting point and test temperature increased. Peterson (92) also pointed out that these compounds, which lubricate fairly well at high temperature, were not effective at room temperature. He explained this difference by stating that both friction and heat are components of the total energy required to break bonds in shear. At low temperatures, more frictional

energy is required; as temperatures approach a solid's melting point, more heat is supplied and less frictional energy is required. Frictional energy is dissipated as heat, which is manifested by the asperity contact temperature rise or "flash temperatures".

ASPERITY CONTACT TEMPERATURES FOR MoS₂-Sb₂O₃ COMPACTS

The hypothesis developed herein for bulk additions to solid lubricants may be plausible if it is possible to demonstrate that the flash temperature is sufficient to soften or deform the bulk additive, resulting in densification of the compact and retention of an overall improved tribological orientation of the primary solid lubricant.

Flash temperatures may be calculated using the notation of Rabinowicz⁽⁹³⁾ and the methodology of Archard.⁽⁹⁴⁾ However, before a flash temperature calculation can be made, one must determine the size of an asperity contact. In the experiments of Quinn and Winer,⁽⁹⁰⁾ hot spots were approximately $50-100~\mu m$ in size, and the number observed at any one time found by successive shortening of photographic exposure time was 7 ± 3 . It was also estimated that the ratio of total hot spot area to apparent contact area was 1:1000.

For the ${\rm MoS}_2{\rm -Sb}_2{\rm O}_3$ tests, the worn pin diameter at test completion is on the order of one millimeter, and if one assumes that the dominant interfacial material is ${\rm MoS}_2$.

then the hardness is 1.5 Mohs for ${\tt MoS}_2$ which is approximately equivalent to 30 kg/mm² on the Vickers scale. Then

$$A_r = \frac{0.1 \text{kg}}{30 \text{ kg/mm}^2} \text{(pin load)}$$

$$= 0.00333 \text{ mm}^2$$
(6.1)

where A_r is the real contact area. Since A_c is the apparent contact area = $\pi r^2 = \pi (0.5 mm)^2 = 0.7854 mm^2$, $\frac{A_r}{A_c} \approx 0.0042$ or $\approx 4:1000$

To which is of the correct magnitude compared to Quiun and Winer's data.

Consider two cases, probably the worst cases: Case A; the entire real area of contact is concentrated in one asperity, and Case B; the real area is split into three equal area contact zones that determine a plane. For Case A, the contact area equals A_r . The contact radius will be

$$r = (\frac{0.00333}{\pi})^{1/2} = 0.0326 \text{ mm} = 3.26 \times 10^{-5} \text{m} \approx 35 \mu\text{m}$$
 (6.2)

or, approximately 70 µm for asperity contact or hot spot diameter. The calculated diameter is within the same size range as those observed by Quinn and Winer, so it is reasonable to assume that only one asperity is in contact

at some times. For Case B, each contact will have an area of

$$\frac{0.00333 \text{ mm}^2}{3} = 0.00111 \text{ mm}^2 \tag{6.3}$$

and the asperity contact or hot spot radius is,

$$r = \left(\frac{A}{\pi}\right)^{1/2} = \left(\frac{0.00111 \text{ mm}^2}{\pi}\right)^{1/2} = 0.0188 \text{ mm} = 1.88 \times 10^{-5} \text{ m}$$

or, approximately 40 µm for asperity contact or hot spot diameter. The calculated diameter is somewhat smaller than the diameter of asperity contacts observed by Quinn and Winer but very near their low value which indicates that the estimate is reasonable.

To calculate the flash temperature, θ_m , Rabinowicz⁽⁹³⁾ provides two equations derived from Archard.⁽⁹⁴⁾ At low speeds,

$$\theta_{m} = \frac{\mu L v}{4 J r (k_1 + k_2)} \tag{6.5}$$

At high speeds,

$$\theta_{m} = \frac{\mu L v^{1/2}}{3.6 L (n c r^{3} k)^{1/2}}$$
 (6.6)

where

µ = friction coefficient

L = load (Newtons)

v = sliding velocity (meters/second)

J = mechanical equivalent of heat (J/cal)

k₁, k₂ = thermal conductivities of contacting materials

p = density

c - specific heat

In order to choose the appropriate equation, Rabinowicz⁽⁹³⁾ supplies an Archard-derived dimensionless parameter⁽⁹⁴⁾ to determine which equation to use. The parameter closely resembles the Peclet number:⁽⁹⁵⁾

$$\lambda = \frac{\text{vrpc}}{2k} \tag{6.7}$$

Rules of application are: if $\lambda < 0.1$, use Equation 6.5; if $\lambda > 5$, use Equation 6.6; if $0.1 < \lambda < 5$, use an intermediate value. In this case, λ is difficult to determine. Archard mentions that thin films on the substrate material (extended surface) will not influence the flash temperature calculation if its thickness is larger than, or much, such smaller than the asperity contact size. In this case, however, the thickness of the surface film is smaller than the asperity contact size, but of the same magnitude. Therefore, some allowance must be

made for both surface film and substrate thermal effects. Archard also defines λ as

$$\lambda = \frac{\tau_1}{\tau_2} \tag{6.8}$$

where τ_1 is the time it takes for heat generated on the surface of contact to reach a point underneath the contact area of depth r, and τ_2 is the time it takes for the asperity to travel through a distance r. If the surface film has thickness t, then the time it takes for surface heat to reach "r" meters beneath the surface becomes:

$$\tau_1' = \frac{t^2 \rho_3 c_3}{2k_3} + \frac{(r-t)^2 \rho c}{2k}$$
 (6.9)

where subscript 3 refers to properties of the surface film. Therefore, if $\tau_2 = \frac{\Gamma}{V}$, then combining Equations 6.7, 6.8 and 6.9, one has

$$\lambda = \frac{\frac{t^2 \rho_3^c}{2k_3} + \frac{(r-t)^2 \rho c}{2k}}{\frac{r}{v}} = \frac{v}{2rkk_3} \left[k \rho_3 c_3 t^2 + k_3 \rho c (r-t)^2 \right] \quad (6.10)$$

In this case, $v = 0.62 \frac{m}{s}$ and $t = 10 \mu m = 10^{-5} m$. Other values were obtained from The Chemical Engineer's

Handbook, (95) The Handbook of Chemistry and Physics, (28)

and a report by Harmer and Pantano. (8)

Por stainless steel:

$$k = 26 \text{ J/m-s-}^{\circ}\text{C}$$

$$c = 460 \text{ J/kg-}^{\circ}\text{C}$$

$$\rho = 7767 \, kg/m^3$$

For MoS₂/Sb₂O₃:

$$p = 3320 \text{ kg/m}^3$$

For MoS_2 , c = 19.7 + 0.00315 T (T in K, c in cal/mol-K)

For Sb₂O₃, c = 19.1 + 0.0171 (600) = 29.4 cal/mol-K = 100.9 cal/kg-*C. Assuming equal contributions to specific heat based upon molar percentage:

$$C_{compact} = (0.75 \times 134.9) + (0.25 \times 100.9) = 126.4$$
 $cal/kg^{\circ}C \times 4.19 \text{ J/cal} = 529.6 \text{ J/kg}^{\circ}C$
(6.11)

Flash temperature calculation: (No units given - all SI units used)

$$\lambda = \frac{(0.62)}{2(3.26 \times 10^{-5})(26)(0.0519)}$$

$$x[(26)(3320)(529.6)(10^{-5})^{2}$$

+ $(0.0519)(7767)(460)(3.26 \times 10^{-5} - 10^{-5})^{2}$ (6.12)

$$\lambda = 32.6 \tag{6.13}$$

Using Equation 6.6, with μ = 0.1, L = 0.98N, J = 1 and all other values given

$$\theta_{\rm m} = \frac{(0.1)(0.98)(0.62)^{1/2}}{(3.6)(1)[(3320)(529.6)(3.26 \times 10^{-5})^3(0.0519)]^{1/2}}(6.14)$$

$$\theta_{\rm m} = 381\,^{\circ}\text{C} \tag{6.15}$$

Values for $\mathfrak c$, c and k are for the $\mathsf{MoS}_2\mathsf{-Sb}_2\mathsf{O}_3$ compact, since it is the surface in contact.

Therefore, at 316°C operating temperature, contact temperature = 316°C + 381°C = 697°C

$$\lambda = \frac{(0.62)}{2(1.88 \times 10^{-5})(26)(0.0519)}$$

$$\{(26)(3320)(529.6)10^{-5})^{\frac{1}{2}}$$

+ $(0.0519)(7767)(460)(1.88 \times 10^{5}-10^{5})^{\frac{1}{2}}$ (6.16)

$$\lambda = 56.0 \tag{6.17}$$

again, using Equation 6.6, as above, except $L = 0.327N_c r = 1.88 \times 10^{-5}m$

$$A_{m} = \frac{(0.1)(0.327N)(0.62)^{1/2}}{(3.6)(1)[(3320)(529.6)(1.88 \times 10^{-5})^{3}(0.0519)^{1/2}}$$
(6.18)

$$\theta_{m} = 290 \, ^{\circ} C$$
 (6.19)

Therefore, at 316°C operating temperature, contact temperature = 316°C + 290°C = 506°C.

These calculations indicate that the flash temperatures are high enough to cause localized softening of the $\mathrm{Sb}_2\mathrm{O}_3$, which has a melting point of 656°C. Softening implies lower shear strength, i.e., less resistance to flow, which enhances the effectiveness of the solid as a lubricant. Ernst and Merchant (96) developed a relationship, incorrectly reported by Peterson, et al. (91),

that the shear strength of a crystalline solid is an exponential function of its melting point.

Their equation is,

$$S = 0.142 \text{ Lpln}(\frac{T_m}{T})$$
 (6.20)

when S = Static shear strength of crystalline solid (kg/mm²)

L = Latent heat of fusion (cal/g)

 ρ = Density (g/cm³)

T = Absolute test temperature (K)

 $T_m = Absolute fusion temperature (K)$

Thus, temperature has a strong influence on the shear strength of an additive. This, along with the analysis of asperity contact temperature lends support to the additive hypothesis.

If the role of $\mathrm{Sb}_2\mathrm{O}_3$ is to soften at asperity contact temperatures, then other materials may perform in a similar fashion. Other materials would cause flash temperatures to be different because of their different material properties. Therefore, the optimum operating temperatures of solid lubricant compacts containing other materials will probably be different from that of $\mathrm{MoS}_2\mathrm{-Sb}_2\mathrm{O}_3$ compacts.

The optimum operating temperature also changes if load or sliding velocity are changed. Once material selection

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has been made, these are the only factors in the flash temperature calculations over which one has some control, as seen in Equation 6.6. If the load is changed, the flash temperature changes proportionately. If the sliding velocity is changed, the flash temperature changes parabolicially. Therefore, since load and speed influence the flash temperature of sliding pairs, these factors would also influence the optimum ambient temperature for a solid lubricant compact. Conversely, load and speed should be optimized for a given operating temperature of a particular sliding pair.

Based upon this analysis, one can predict the temperature range in which the additive will be most effective for a given set of conditions. Once materials are selected, the load and speed conditions must be specified in order to determine the flash temperature. The asperity contact temperature should be in a range that permits the additive to effectively deform. At temperatures too high, the additive will shear too easily, while at very low temperatures the additive rannot easily shear so that beneficial deformation does not occur.

CHAPTER VII

OTHER OXIDES AND SULFIDES AS BULK ADDITIONS TO MoS2

WEAR STUDIES

Based upon the development of an hypothesis for bulk additions in MoS_2 , several oxides and sulfides were selected as additives for MoS_2 to determine if tribological performance would be as predicted by the hypothesis.

Additives were selected primarily on the basis of earlier efforts by many investigators to find stable materials that would improve the performance of a variety of solid lubricants. Additives of this type, which typically have low values of Mohs hardness include As₂O₃, Bi₂O₃, Sb₂S₃, Sb₂S₄, PbO, CdO, and CuO. Other additives were selected because of their high hardness, i.e., TiO₂ and Al₂O₃, so that a comparison with softer materials could be made. Additionally, MoO₃ was selected since it might be an oxidation product of MoS₂ found at the interface.

All of these additives were ground, with the exception of PbO, on the machanical amalgamator for a period of 16 minutes followed by sieving through a 30 µm nickel screen. PbO only sieved prior to compaction with MoS₂ since it oxidizes readily in the amalgamator, easily observed because of the color change.

Each of the sieved oxides or sulfides was weighed to the nearest tenth of a milligram and formulated based upon the volume equivalent to that of $\mathrm{Sb}_2\mathrm{O}_3$ additions to MoS_2 . An exception was MoO_3 which was added on a molar basis for comparisons with possible eutectic compositions of $\mathrm{Sb}_2\mathrm{O}_3$ and the oxidation products of MoS_2 .

With few exceptions, only a single compact of MoS_2 and each oxide or sulfide was prepared in a procedure identical to the one used for $MoS_2-Sb_2O_3$ compacts. Since these experiments were designed only to establish trends in wear volume and friction coefficient as functions of additive properties, repeat evaluations were not performed.

The compacts were shaped and then worn at 316° C on stainless steel shim stock as done with $805_2-80_20_3$ compacts. The data generated as given in Appendix D are summarized in Table 20 along with the Mohs hardness (97) and temperature of melting, transition, sublimetion or decomposition (28) for each additive.

As seen in Table 20, the variation in wear volumes was large, varying from 0.04 mm^3 to more than 0.4 mm^3 . Coefficients of friction were as low as 0.06 and as high as 0.24 during wear tests. The lowest wear volumes were found for $\mathrm{Sb}_2\mathrm{S}_4$ and MoO_3 additions. These wear volumes were approximately one-half of the wear generated when $\mathrm{Sb}_2\mathrm{O}_3$ was the additive in MoS_2 . These additives were beneficial if the Mohs hardness was less than or equal to 4.5. TiO_2 and

Table 20. PELLET WEAR IN DRY AIR AT 316°C

PELLET COMPOSITION	TEST	MELTING POINT* (°C)	MOHS HARDNESS	WEAR VOLUME (mm ³)	F _K	3. 14.	la La
MoS ₂		1185	1-1.5	0.157	0.07	0.08	0.68
Mos ₂ + Additive							
A1,01	146	2072	•	0.415	0.12	0.24	0.17
A8203	145	312 (8)	1.5	0.070	0.11	0.16	0.14
B1203	132	825	4.5	960.0	0.14	0.23	0.19
c q o	133	(P) 006	E	0.071	0.10	0.17	0.14
Ono .	136	1326	3.5	0.072	90.0	0.16	0.11
Hoo,		795	7	0.044	0.11	0.17	91.0
Pbo	130	886	7	0.061	0.12	0.20	0.16
T10,	135	1840	9	0.432	0.14	0.21	0.18
Sboog		959	2.75	0.093	0.09	0.14	0.12
Sb_03		550	7	0.088	0.11	0.17	0.13
Sb2S4	134	210	ю	0.043	0.14	0.29	0.17

s = sublines; d = decomposes

 ${\rm Al}_2{\rm O}_3$ produced greater wear volume than did ${\rm MoS}_2$ without additives. Those additions to ${\rm MoS}_2$ produced wear volumes that were about two end one-half times greater than the wear volume for ${\rm MoS}_2$, which may result from the inability of those oxides to deform under the tribological conditions of the test.

In Figure 36, wear volume is plotted as a function of Mohs hardness. The least square regression equation is as follows:

Wear Volume $(mm^3) = 0.0563$ (Mohs Hardness) - 0.0664 (7.1) Correlation coefficient r = 0.88

Therefore, about 78 percent of the variation in wear volume may be attributed to differences in hardness of the additive.

When the linear correlation coefficient is calculated for wear volume as a function of the absolute melting temperature of the additive, a result comparable with that of wear volume as a function of hardness. The linear regression equation is:

Wear Volume
$$(mm^3)$$
 = 2.23 x 10^{-4} (Melt. Temp., K) = 0.0774 (7.2) r = 0.87

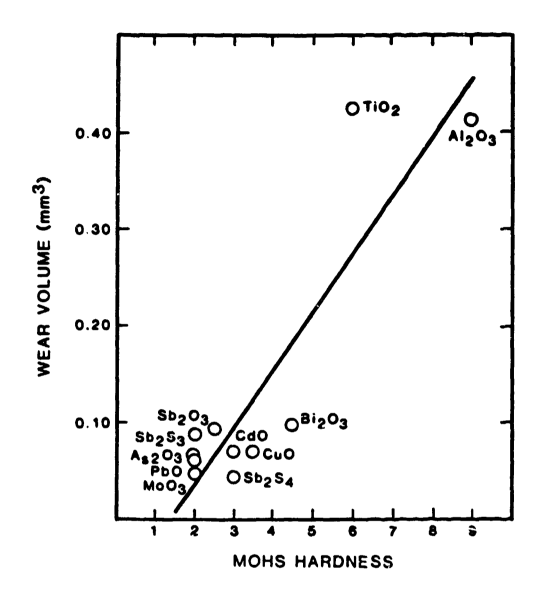


Figure 36. Wear Volume as a Function of Mohe Hardness.

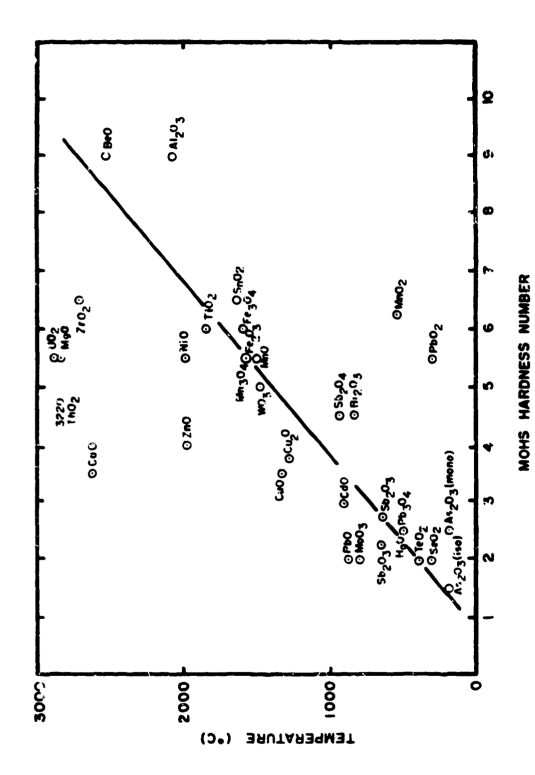
Such a correlation is expected since Mohs hardness varies linearly, with some exceptions, with the melting temperature of materials. (85,89) For oxides, such a relationship is presented in Figure 37.

Correlation of initial coefficient of friction (30 sec), μ_{k_1} , or final coefficient of friction (15 min), μ_{k_1} , is with Mohs hardness or melting temperature is poorer. Linear regression suggests that correlation of basic additive properties is better with final coefficient of friction values (r = 0.66 for correlation with Mohs hardness, r = 0.59 for correlation with melting temperature) rather than with initial friction coefficient values (r = 0.30 and r = 0.295).

The data suggest that it is difficult to predict $\mu_{k_i} \text{ or } \mu_{k_f} \text{ by use of Mohs hardness or melting } \\ \text{temperature. However, as seen in Figure 38, some trending of initial, mean and final friction coefficient as a function of Mohs hardness is apparent as reflected an higher correlation coefficients as frictional coefficients increase during the test.$

Thus, it appears that the wear volume generated is related to the hardness and melting temperature of the additive, but the friction coefficient is not.

Mohs hardness reflects the shear strength of the additives. However, Mohs hardness is measured at ambient and wear volume was determined at 589 K with asperity



Melt Temperature of Oxides as Function of Hohe Hardness. Figure 37.

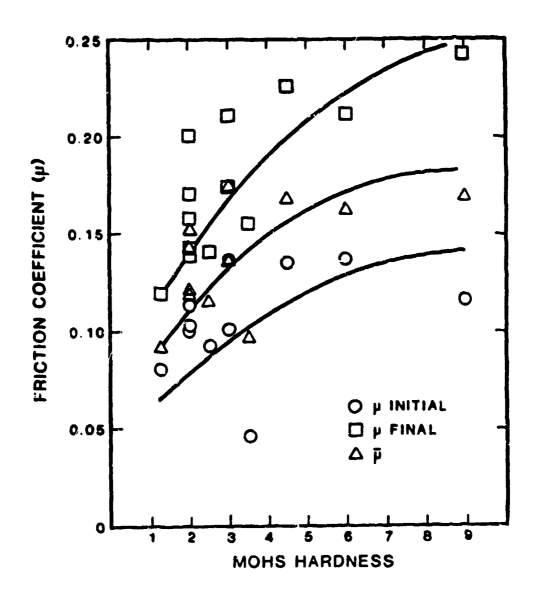


Figure 38. Friction Coefficient of ${\rm MoS}_2{\rm -Oxide}$ Compacts as a Function of Mohs Hardness of Oxides.

contact temperatures even higher. Therefore, correlations of a) wear volume as a function of the difference between softening point and test temperatures, and b) wear volume as a function of additive shear strength at test temperature were calculated.

Handbook data for the required quantities and calculated values to be employed later are compiled in Table 21.

The relationship relating wear volume and friction coefficient to the difference of G.? absolute melting temperature and the test temperature was evaluated initially. 0.7 of the melting temperature is at the upper limit for transition from brittle to ductile behavior for materials similar to oxides and sulfides evaluated here. (98) That transition is, of course, related to deformation of the additive as hypothesized. The linear regression equation for wear volume is as follows:

Wear Volume
$$(\pi m^3) = 3.18 \times 10^{-4} (0.7 T_m^{-589}) + 0.0492$$
(7.3)

r = 0.87

The correlation coefficient suggests that wear volume is correlatable with the difference of 0.7 T_m -589. Attempts to correlate friction coefficient with 0.7 T_m -589 resulted in correlation coefficients of 0.20 (initial) and

Table 21. ADDITIVE DATA FOR EVALUATION OF SHEAR STRENGTH AT TEMPERATURE

ADDITIVE	ADDITIVE MEAT OF FUSION DENSITY	DE#8114	HELT TEMPERATURE 0.7 To-369 To	9.7 T ₆ -569	-° -	8 14388	SHEAL STRENGTU WPs	VEAR VOL(ma ³)	مو	بمو
	(8/170)	(8,00)	7. (E)			bg/m² (8	kg/mo ² (H/m ² = 10 ⁻⁶)			•
1,20,	1	3	8 €.01	179.4	:	11.49	112.66	9.00	4.134	
10201		2.67	929	•1.3	1.577		164.52	6.693	•	•
2		9.53	1159	271.3	1,96	11.54	113.17		• 113	
Ne0,		£.69	••01	138.0		•	67.20	0.014	:	=
0.0		•	1999	556.3	2.715	32.14	315.20	6.072	•	•
070		6.15	1773		3.010	:				
1102		4.26	2113	1.06	3.567	169.2	1659.30	0.413	9.137	
41203	256	3.965	2345	1052.5	3.983	139.1	1952.60	6.415	•: ::	0.24
4020)		4.15	\$88	-179.5		-0.092	-0.90(•)	0.07	:	•
		•	623	- 12.9	1.397	1.27	71.39	0.000		0.17
St. 2 54										
TW SI.	•									

0.59 (final), which suggest that friction coefficients are not strongly related to 0.7 $T_{\rm m}$ -589.

Computed negative values of 0.7 T_m -589 suggest that the softening temperature of the additive is exceeded in tests based upon that relationship. If those values are excluded from consideration, the linear regression equation relating wear volume to 0.7 T_m -589 is as follows:

Wear Volume
$$(mm^3) = 3.928 \times 10^{-4} (0.7 T_m - 589) - 0.0016$$
(7.4)

r = 0.91

Therefore, the correlation is improved when negative values of (0.7 $T_{\rm m}$ =589) are eliminated.

For an estimate of the shear strength of additives at test temperature, the theoretical relationship of Ernst and Merchant $^{(96)}$ as given in Equation (6.2) was employed. Values of the shear strength of the additives were not calculated for CdO and $\mathrm{Sb}_2\mathrm{S}_4$ since pertinent data were not available. The linear regression equation relating wear volume to shear strength as calculated is as follows:

Wear Volume
$$(mm^3) = 0.0002 (N/m^2 \times 10^{-6})$$

+ 0.0524

r = 0.983 (7.5)

መጀታሪያ የለፈራጊ እና የሚያስፈርር እና የሚያስ

Wear volume data plotted as a function of shear strength is presented in Figure 39. Linear correlation coefficients for μ_k and μ_k as functions of calculated shear strength are 0.37 and 0.62, respectively, which are comparable with the correlation of μ with additive hardness and melting temperature. Thus, the wear volume is related very strongly to the calculated shear strength. Initial and final coefficients of friction are much more weakly related to shear strength. The high correlation coefficient between wear volume and calculated shear strength of the additive indicates that more than 97 percent of the variation in wear volume may be attributed to differences in calculated shear strength at test temperature.

The weakness of correlation of the friction coefficient shear strength indicates that the beneficial effect of an additive is not an interfacial phenomenon. Therefore, the effect must occur within the bulk of the solid lubricant, which supports the hypothesis of additive deforming to permit the solid lubricant to attain and retain a preferred orientation for improved tribological condition.

The weaker correlation of wear volume with additive hardness, melting temperature, or 0.7 T_m -589 might be expected sine; they are related to shear strength of the additives, but are, at best, poor predictors of shear strength at test temperature.

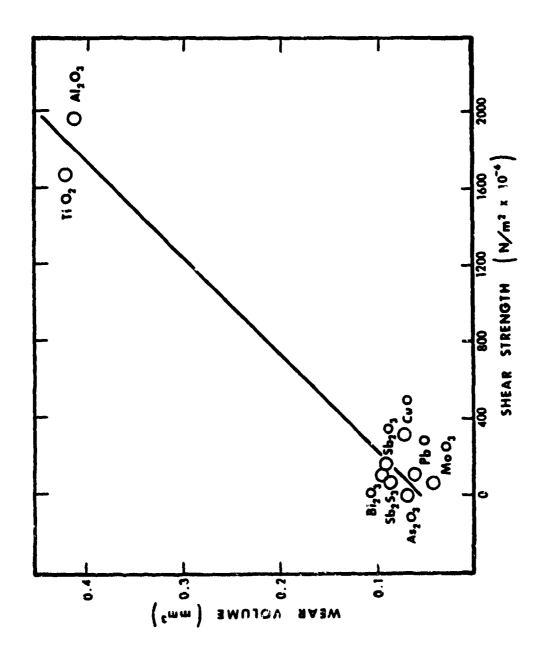


Figure 39. Wear Volume as Function of Additive Shear Strength.

Sb204 AS A BULK ADDITIVE

Wear of compacts of MoS_2 containing Sb_2O_4 were of special interest, since it may be possible for Sb_2O_3 to oxidize to Sb_2O_4 resulting in different tribological properties. Data are presented in Table 22 summarizing the results of the assessment. The wear of such compacts were higher than similar compacts containing Sb_2O_3 (o) (Tests 84-86). These data indicate that oxidation of Sb_2O_3 to Sb_2O_4 at the compact interface would not be beneficial.

Moo3 and Moo3-Sb2O3 AS BULK ADDITIVES

 ${
m MoO_3}$ as an additive and in combination with ${
m Sb_2O_3}$ was evaluated to examine whether any synergism or eutectic formation occurs between it and ${
m Sb_2O_3}$.

MoO₃ has some capability as a solid lubricant. It has been documented that molybdenum sliding against molybdenum in air under mild sliding conditions has rather high friction (µ-1) up to about 480°C, after which the friction remains low (µ=0.3) until about 790°C. (99) These temperatures correspond quite well with the onset of rapid molybdenum oxidation to MoO₃ and with melting point of MoO₃, respectively. As temperature is decreased below 480°C, friction is lower on MoO₃ than the bare metal, but higher than at temperatures from 480°-790°C. Under the conditions of the test, there is, apparently, about a 300°C temperature range below the melting point of MoO₃ in which it acts as a fair solid lubricant.

Table 22. EFFECT OF TEST TIME ON WEAR VOLUME OF ${\sf MoS_2\text{-}Sb_2O_4}$ COMPACTS

(DRY AIR, 316°C; 100 g LOAD)

TEST Number		DITIVE PERCENT	TE		R TIME 1E (mir	n.)
			1.0	2.5	5.0	7.5
102	25	sb ₂ 0 ₄	0.077			
100	25	Sb204		0.099)	
98	25	Sb ₂ 0 ₄			0.108	3
101	25	sb ₂ 0 ₄			0.150)
99	25	Sb ₂ G ₄				0.167

Data generated in wear evaluations of MoO_3 , Sb_2O_3 , and $MoO_3-Sb_2O_3$ in MoS_2 compacts are summarized in Tables 23, 24, 25, 26 and 27.

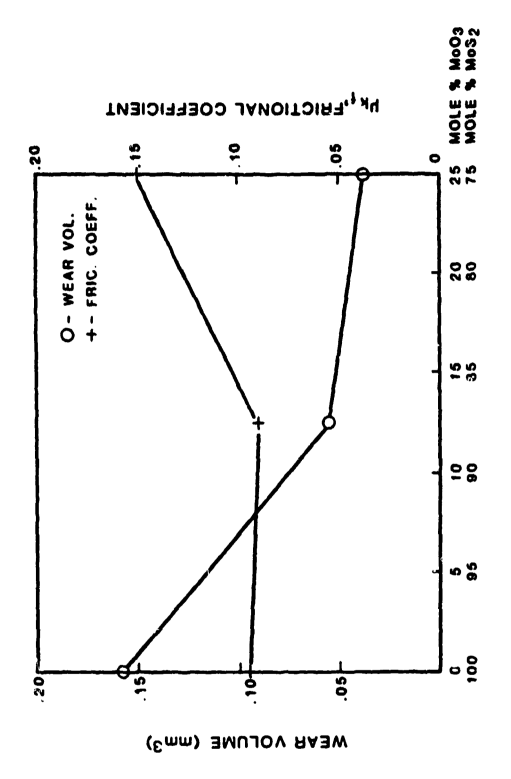
Wear volumes and friction coefficients are plotted as functions of composition for MoO_3 in MoS_2 , respectively, in Figure 40.

These data indicate, contrary to an assumption that MoO_3 could not be a beneficial additive because of its abrasive nature, (99,100) wear compacts containing MoO_3 produced only about half the wear volume of MoS_2 -Sb $_2O_3$ compacts.

In Figure 41, the wear volumes of ${\rm MoO_3}$ and ${\rm Sb_2O_3}$ in ${\rm MoS_2}$ are compared on a molar basis. It is seen that ${\rm MoO_3}$ is superior in wear volume at each concentration.

Finally, Figure 42 shows that a MoS_2 compact containing equal parts of MoO_3 and Sb_2O_3 gives a wear volume between the wear volume of MoS_2-MoO_3 and $MoS_2-Sb_2O_3$ compacts. This observation, combined with the fact reported earlier that a eutectic of $Sb_2O_3-MoO_3$ could not be found, shows that there is no synergism between Sb_2O_3 and MoO_3 . The results do show, however, that mixtures of MoS_2 with MoO_3 can be better solid lubricants than $MoS_2-Sb_2O_3$ under these test conditions.

When present with MoS_2 , it is claimed that MoO_3 aids in forming a continuous film. (82) Peace (65) suggests that the oxide may act as a binder for MoS_2 . Winer (17) suggests



Pigure 40. Wear Volume and Priction Coefficient for MoS₂-Sb₂0₃ Compacts.

Table 23. WEAR VOLUME OF Mos_2 AND Mos_2 WITH Sb_2o_3 , OR Mos_2 WITH Moo_3 AND Sb_2o_3 (DRY AIR, 316°C; 100 g LOAD)

TEST	ADDITIVE	WEAR VOLUME
NUMBER	MOLE PERCENT	TEST TIME (min.)
		1.0 2.5 5.0 7.5 16.7
88		0.047
90		0.111
89		0.120
91		0.128 0.172
92	₩ 🕳 🗢	
84	25 Sb ₂ O ₃ (o)	0.049
8.5	25 Sb ₀ O ₂ (y)	0.060
86	25 Sb ₂ O ₃ (v) 25 Sb ₂ O ₃ (o)	
0.073	2-3	
110	3.75 MoO ₃	0.079
	3.75 Sb ₂ 03(0)	
106	7.5 Mo03	0.087
	7.5 Sb ₂ O ₃ (o)	
109	17.5 Moő3	0.082
	17.5 Sb ₂ O ₃ (o)	
104	10 MoO ₃	0.074
_	5 Sb ₂ O ₃ (o)	
107	20 MoO ₃	0.053
	5 Sb ₂ O ₃ (o)	

Table 24. WEAR VOLUME OF MOS_ WITH ${\rm Sb_2O_3}$, OR MoS_ WITH MoO_3 and ${\rm Sb_2O_3}$ (DRY AIR, 316°C; 100 g load)

MUMBER	ADDITIVE MOLE PERCENT		WEAR TEST TI	WEAR VOLUME Test time (min.)		
		1.0	2.5	2.0	7.5	16.7
8	•	0.047				
90	•		0.111			
68				0.120		
91	1 1			0.128		
92	:				0.172	
8.4		0.049				
8 5	25 Sb ₂ 0 ₃		0.060			
86	25 Sb ₂ 0 ₃ (o)			0.073		
011	3.75 HoG2					0.079
	3.75 Sb, 0, (o)					
106	•					0.087
	•					
109	17.5 Moog					0.082
	17.5 sb ₂ 03(c)					
104	10 MoO.					0.074
107	20 MoO ₃					0.053

WEAR VOLUMES AND PRICTION COEPFICIENT FOR Table 25.

	AND	Mosz, Mos Compositions	HOS2 AND HOO IONS (DRY AIR	3. H	Mos ₂ WITH 589 k, 100	TH MOO3-Sb ₂ O3	5.7	, Î	
TEST NUMBER	ADDITIVE HOLE PERCENT	WEAR (NE ³)			18	TEST TIME (min.)	min.)		
			5.	1.0	2.5	5.0	7.5	0.1	21
					PRIC	PRICTION COE	COEFFICIENT		
114	!	0.181	190.	190.	.061	.061	.061	.072	780
115	•	0.134	.075	.075	.075	.085	.095		
113	5	0.075	• 079	•019	080	.088	960.	100	
9 ::	25 Sb ₂ 0 ₃	0.105	. 109	.108	.108	1111	111.	.113	.115
571	n ,	0.108	820.	*60 •	. 103	. 113	.121	.127	. 136
671	n (0.078	001	. 108	.123	.162	.165	. 165	. 192
711	;	0.057	.072	•076	.076	.079	060.	.097	. 100
9 .	E:	0.037	!	1	!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!	* • •	•	1	!
- :	ž :	0.055	100	. 100	. 100	.115	. 119	. 130	041.
<u> </u>	e F	0.045	.127	.127	.133	.145	.151	.163	.171
e kalif =1 •		0.031	!!!	!	1	!	!		
171	5 HoO3	0.050	• 106	.111	.123	.130	.142	.147	. 1.58
		0.059	!	1	!	;			
104	2.5 Sb	0.074	1	!	:		!	1	;
8 1 18		0.057	.078	180.	*60 *	.103	.112	.119	. 129
122	2.5 Sb2 2.5 Mo3	0.0811	.075	.081	960.	.107	.107	.111	. 136
125	5 Sb ₂	0.079	.086	.095	.110	.129	.144	.154	.178
131	.5 8b2	0.036	.067	.071	.085	.109	.124	.142	.162
138	• 5 Sb ₂	0.077	180.	.084	.089	, 106	.123	.137	.169

le 26. EFFECT OF VARYING MOO3-Sb203 ADDITIVE CONCENTRATION ON PELLET WEAR VOLUME (316°C, 1000 SEC., DRY AIR, 100 g LOAD) Table 26.

McO3: Sb2O3		WEAR V	WEAR VOLUME (mm ³) and (TEST HOS ₂ MOLE PERCENT	and (TEST NO.)	(-	
Molar Ratio	6.5	7.5	85	87.5	92.5	100
	.820 (109)	.059 (103) .057 (1:8) .074 (104) .081 (122) .079 (125)	.086 (106)		.079 (110)	
2:1			.043 (105)			
1:5		.053 (107)				
1:0		.045 (117) .037 (108) .055 (111) .031 (119)	.0572 (112)			
0:1		.078 (124) .108 (123) .105 (116)	.075 (113)			
0:0						.181 (114)

		OF PRICTION	(316°C,	DRY AIR, 10	OF PRICTION (316°C, DRY AIR, 100 g LOAD)	on coefficient	2 2 1
MoO3:5b2O3		COEPPICIE	NT OF FR	COEFFICIENT OF FRICTION (µ) AND (TEST MOS ₂ HOLE PERCENT	IND (TEST NO.)		
	65	7.5	85	87.5	92.5	100	
1:4		.1314					
1:1	1	.0813 .0507 .0814	!		•		
2:1							
4:1		:					
1:0		.1117		.0710			
0:1		.0814		.0811			
0:0						6090	

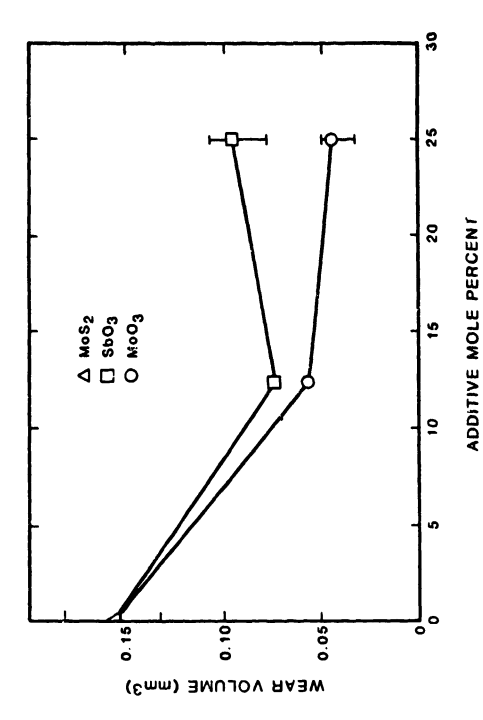


Figure 41. Wear Volume for MoO_3 - MoS_2 and Sb_2O_3 - MoS_2 Compacts.

K Systyl miteratoroum selectoroum ponerentorm ponerent

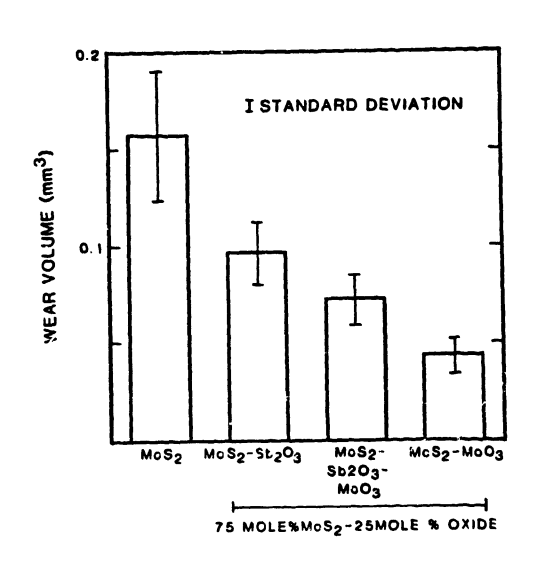


Figure 42. Wear Volume for MoS2-NoO3-Sb2O3 Compacts.

that the oxide formed on the outer layer of MoS_2 reduces the oxidation of the underlying layers of MoS_2 . Thus, low values of friction are maintained in a MoS_2/MoO_3 mixture as long as a subfilm of MoS_2 remains. Fleischauer and Bauer's (101) results show that for long wear lifetimes, 30 to 40 percent of the interfacial layer for MoS_2 should be oxidized.

The MoS₂-MoO₃ formulation in this effort was not optimized tribologically; however, the concentration of MoO₃ in MoS₂ was, coincidentally, near the optimal concentration determined by Fleischauer and Bauer. This work and the observations of others support the hypothesis advanced that oxides, such as MoO₃ or Sb₂O₃, which can deform under environmental conditions encountered, aid in establishing a tribologically superior interface established by retaining the solid lubricant in referred orientation.

The tests with oxides other than $\mathrm{Sb}_2\mathrm{O}_3$, especially MoO_3 , prove that $\mathrm{Sb}_2\mathrm{O}_3$ is not unique in its beneficial role in MoS_2 and that a eutectic of $\mathrm{MoO}_3\mathrm{-Sb}_2\mathrm{O}_3$ that could be expected to form for compacts of $\mathrm{MoO}_3\mathrm{-Sb}_2\mathrm{O}_3$ composition tested, do not play a significant tribological role.

CHAPTER VIII

GRAPHITE LUBRICANT COMPACT SYSTEM

In addition to MoS₂, graphite is often employed as a solid lubricant. In contrast to MoS₂ graphite requires chemisorption of oxygen at surface cracks, physical adsorption of water vapor, or some similar mechanism to reduce cleavage strength for good tribological performance. (102) If used in air, the performance of graphite rapidly decreases as a function of increasing temperature up to about 200°C because it desorbs such materials. At much higher temperatures, i.e., 425°C and above, performance again improves (84) possibly resulting from a more active role for oxygen.

ASPERITY CONTACT TEMPERATURE

To examine systems other than those using MoS_2 as the solid lubricant, an investigation of graphite containing antimony trioxide was conducted. Conditions in the graphite $8b_20_3$ system compared with $MoS_2-Sb_20_3$ suggested that Sb_20_3 might deform at the asperity contacts at different temperatures. A critical factor is the higher friction coefficient of graphite at the test temperature.

Other properties were also predicted to affect flash temperatures.

To estimate asperity contact flash temperatures, the procedure of Archard $^{(94)}$ was followed. First, the real area of contact is calculated.

$$A_{r} = \frac{L}{P} \tag{8.1}$$

where

 A_r = area of real contact

L = load

P = hardness

Since the hardness of graphite is about 0.5 on the Mohs scale, this corresponds to about 63% less than MoS_2 hardness as shown by Tabor. (87) Thus,

$$P_{graphite} = 0.63 P_{MoS_2}$$
 and since, $P_{MoS_2} = 30 kg/mm^2$ (8.2)

$$P_{graphite} = 0.63 (30 \text{ kg/mm}^2) = 19 \text{ kg/mm}^2$$
 (8.3)

and,

$$A_{r} = \frac{L}{P} = \frac{0.1 \text{ kg}}{19 \text{ kg/mm}^2} = 0.00526 \text{ mm}^2$$
 (8.4)

Two cases as stated in the analysis of the $MoS_2-Sb_2O_3$ compact will be considered.

Case A for single point contact:

$$A_r = \pi r^2$$
 or $r = (\frac{A_r}{\pi})^{1/2}$ (8.5)

$$r = (\frac{0.00526 \text{ mm}^2}{\pi}) = 0.0490 \text{ mm} = 4.09 \times 10^{-5} \text{m}$$
 (8.6)

Case B for three point contact:

$$\frac{A_r}{3} = \frac{0.00526 \text{ mm}^2}{3} = 0.00175 \text{ mm}^2$$
 (8.7)

$$r = (\frac{0.00175 \text{ mm}^2}{\pi})^{1/2} = 0.0236 \text{ mm} = 2.36 \times 10^{-5} \text{m}$$
 (8.8)

The value of λ must be determined using Equation (6.7) to determine which flash temperature equation should be used for the final calculation. Once again v = 0.62 m/s and $t = u = 10^{-5}$

For stainless steel:

$$k = 26 J/m-s-C$$

$$c = 460 \text{ J/kg} - {}^{\circ}\text{C}$$

$$\rho = 7767 \text{ kg/m}^3$$

For graphite - Sb₂0₃:

$$k = 0.13 \text{ J/m-s-}^{\circ}\text{C}$$

$$c = 1133 \text{ J/kg-}^{\circ}\text{C}$$

$$p = 2239 \text{ kg/m}^3$$

For graphite:

 $= 3.92 \text{ cal/mol K} = 327 \text{ cal/kg-}^{\circ}\text{C}$

For Sb₂O₃:

$$\rho = 5435 \text{ kg/m}^3$$

c = 100.9 cal/kg-C

Assuming equal contribution according to the law of mixtures for specific heat and density according to pellet composition percentage:

Theoretical density = ρ_{T} = (0.75 x 2169) + (0.25 x 5435) = 2986 kg/m³

For compacted particles, real density is estimated at about 75 percent of theoretical so

$$\rho = 0.75 \text{ x (theoretical density)} = 0.75 \text{ x } 2986 = 2239 \text{ kg/m}^3$$
 $c = 0.75 \text{ x } 327 + 0.25 \text{ x } 100.9 = 270.5 \text{ cal/kg°C} = 1133$
 $J/kg°C$

Therefore:

Case A for single point contact:

$$\lambda = \frac{(0.62)}{2(4.09 \times 10^{-5})(26)(0.18)} [(26)(2239)(1133)(10^{-5}))^{2} + (0.18)(7767)(460)(409 \times 10^{-5} - 10^{-5})^{2}]$$

$$= 11.7$$
(8.9)

Using Equation (6.6), with μ = 0.4, L = 0.98N, J = 1, all other values givan.

$$\theta_{m} = \frac{(0.4)(0.98)(0.62)^{1/2}}{(3.6)(1)[(2239)(1133)(4.09 \times 10^{-5})^{3}(0.18)]}$$
= 485°C (8.10)

Therefore, at 316°C test temperature, contact temperature = 316°C + 485°C = 801°C

Case B for three point contact:

$$\lambda = \frac{0.62}{2(2.36 \times 10^{-5})(26)(0.18)} \times \\ [(26)(2239)(1133)(10^{-5})^{2}+(0.18)(7767)(460) \times \\ ((2.36\times10^{-5})-10^{-5})^{2})]$$

$$= 18$$
(3.11)

Using Equation (6.6), as above, except L = 0.327N

$$\theta_{m} = \frac{(0.4)(0.327)(0.63)^{1/2}}{(3.6)(10[(2239)(1133)(2.36 \times 10^{-5})^{3}(0.18)]^{1/2}}$$

$$\theta_{m} = 369 \, ^{\circ}C \qquad (8.12)$$

therefore, at 316°C test temperature, contact temperature = 316°C + 369°C = 685°C.

Both of the above temperatures are much higher than the melting point of Sb_2O_3 (656°C). To get the same effect as that that occurs in the MoS_2 - Sb_2O_3 system, all other parameters being equal, the ambient temperatures would need to be reduced as follows:

Case A for single point contact:

Contact temperature in $MoS_2-Sb_2O_3$ system = $697^{\circ}C$ Contact temperature in Graphite- Sb_2O_3 system = $801^{\circ}C$ Test temperature reduction necessary = $801^{\circ}C$ - $697^{\circ}C$ = $104^{\circ}C$

316°C - 104°C = 212°C for equivalent test temperature

Case B for three point contact:

Contact temperature in MoS₂-Sb₂O₃ system = 606°C

Contact temperature in Graphite-Sb₂O₃ system = 685°C

Test temperature reduction necessary = 685°C - 606°C = 79°C

316°C - 79°C = 237°C for equivalent test temperature

Therefore, in the standard test, if one reduces test temperatures to about $210^{\circ}\text{C}-230^{\circ}\text{C}$ ($410^{\circ}-446^{\circ}\text{P}$), then asperity contact temperature in the graphite-Sb₂O₃ system

would approximate those in the $MoS_2-Sb_2O_3$ system evaluated at 316°C (600°F).

WEAR STUDIES

To determine if the relative sliding wear of the graphite-Sb₂O₃ system compared with graphite would be improved by lowering the test temperature appropriately, pellets of graphite (U.S. Army Specification, 2-64A, Graphite Powder, Dixon Jersey City, NJ) and graphite-Sb₂O₃ were prepared in the standard configuration and evaluated at 232°C (450°F) and 361°C (600°F). It was predicted as shown earlier that at the higher temperature the matrix of Sb₂O₃ would be expected to deform severely because of localized melting, while at lower temperature the Sb₂O₃ matrix would only soften to accommodate a preferred reorienfation of graphite.

The raw and reduced data are summarized in Table 28. The significance of the teste is that at the higher temperature, the wear rate of the graphite-Sb₂O₃ pellet was much greater than that of graphite due most probably to excessive softening or melting of Sb₂O₃. At the lower temperature of 232°C, the graphite-Sb₂O₃ pellet wore less than a graphite pellet at the same temperature. Here Sb₂O₃ softening rather than melting is expected by the flash temperature calculations.

The friction coefficient data are significant. For graphite, the friction coefficient as a function of time

Table 28. GRAPHITE AND GRAPHITE/Sb203(o) WEAR DATA

1821			7	<u>.</u>	4	41
	i i	retter tonros:1108	WEAR VOLUME (MM)	1 P E E	, r	•
141	21.1	Graphite	0.082	0.10)	0.138	9.123
143	(10)	Graphite/Sb ₂ 0 ₃ (v)	0.1125	0,165	9.1.6	6.170
129	232.2	Graphice	0.0375	0.160	0.271	0.215
128	(450)	Graphite/Sb ₂ O ₃ (o)	0.0342	0.126	0.240	0.183
041	273.9	Graphite	0.0666	0.161	0.198	9.19
142	(525)	Graphits/Sb ₂ O ₃ (o)	0.1246	0.094	0.202	•
127	315.6	Graphite	0.0427	0.227	0.343	0.285
126	(009)	Graphice/8b203	0.0898	0.101	0.247	0.174
		RATIOS OF ME	HORMALIZED DATA			
	21.1	Graphite	1.92	2.47	0.40	0.432
	(10)	Graphite-Sb ₂ O ₃ (o)	2.63	6.73	0.57	0.62
	232.2	Greehite	0.878	0.705	0.190	0.754
	(450)	Graphite-Sb ₂ O ₃ (o)	0.801	0.555	0.100	0.642
	273.9	Graphite	1.419	0.109	0.577	0.632
	(\$2\$)	Graphite-Sb ₂ O ₃ (e)	2.918	0.414	0.589	9.519
	315.6	Graphite	1.00	1.000	1.000	1.000
	(009)	Graphite-5b203(e)	2.10	0.145	0.720	0.611
		CRAPHITE-Sb203(0)/CRAPHITE	RATIO OF	HORANLIZED BATA		
	21.1		1.370	1.553	1.425	1.435
	232.2 (450)		0.912	0.787	9.86	
	273.9 (525)		2.056	0.585	1.02	0.02
	315.6		2.10	0.445	0.726	0.610

was relatively constant, while pellets containing \$b203 produced friction coefficients that increased almost linearly from the initial to final value. At the higher temperature, the initial value, $\mu_{\mathbf{k}}$, was lower than the corresponding value at 232°C while a reversal was observed in final values. These data may be rationalized if SbyOq is severely deforming or melting at the higher temperatures which would contribute to an initial low friction coefficient and deposit material onto the relatively cooler wear track, which would result in a gradual increase in friction coefficient. At lower temperatures, the effect of asperity contact flash temperature on coefficient of friction is minimized. Graphite and graphite-Sb2O3 compacts were also evaluated at room temperature (Tests 141 and 143, respectively) and at 274°C (Tests 140 and 142, respectively). Results at 274°Cwere intermediate between those at 232°C and 310°C, with relative wear of the graphite-Sb2O3 compacts higher than at 232°C. Frictional coefficient ratios were uniformly higher at lower temperatures as shown in Figure 43.

In summary, a predicted by flash temperature calculation, for 274°-316°C test temperatures, additions of Sb₂O₃(o) to graphite are not beneficial. At those temperatures, Sb₂O₃(c) may fuse rather than deform at asperity contact and, thus, the compact wears at a higher rate than graphite alone. When ratios of data for tests

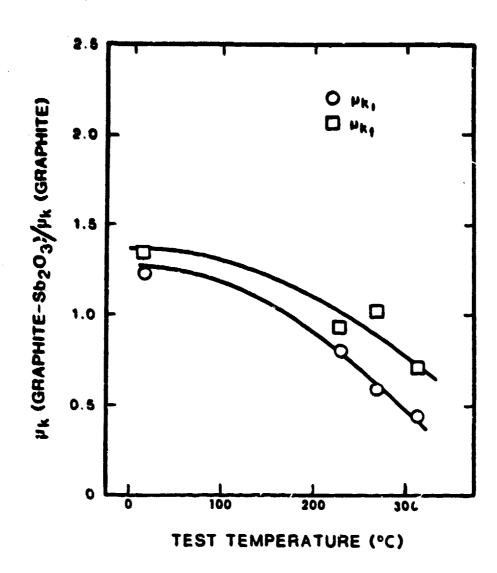


Figure 43. Ratio of Graphite-Sb₂O₃ to Graphite Friction Coefficients as Function of Test Temperature.

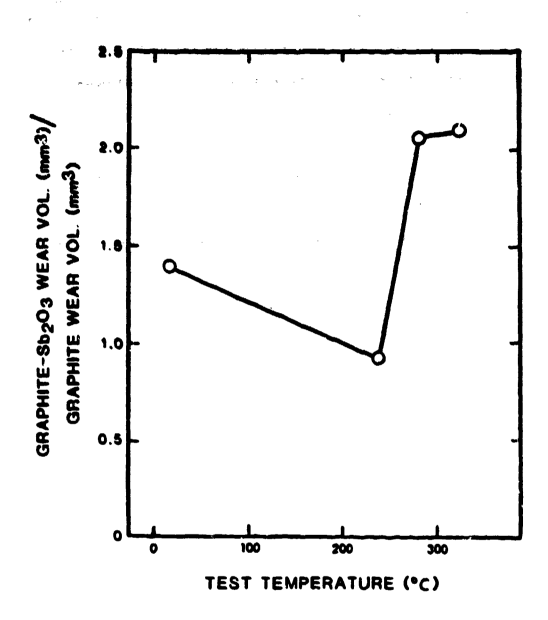


Figure 44. Ratios of Graphite-8b203 to Graphite Compact
Wear Volumes as a Function of Test Temperature.

over the range 21°-316°C are examined as given in Figure 44, it is found that at a test temperature of 232°C, the relative wear volume is less for a compact containing Sb203(0). At ambient, the effect of Sb203 additions to graphite does not appear to be beneficial, which may be due to an inability of the additive to sufficiently deform under those conditions. These data support a concept of bulk additives plastically deforming for improved tribological surfaces under selected conditions. If the asperity contact temperatures are too high, the additive may deform excessively and the addition becomes tribologically detrimental. Of course, other factors such as load and sliding velocity are also crictical factors with temperature in determining the range of conditions over which the additive may be beneficially compounded with a solid lubricant.

The data also indicate that at temperatures in these tests where $\mathrm{Sb}_2\mathrm{O}_2(o)$ might be expected to be beyond its beneficial addition range, i.e., 274° and 316°C, the initial coefficient of friction, μ_{k_1} , is low followed by a gradual increase to a much higher value, μ_{k_1} , by the end of the test. It is hypothesized that the additive may act as a fluid or subline at a very high rate during initial asperity contact; as wear progresses, graphite- $\mathrm{Sb}_2\mathrm{O}_3$ is deposited on the cooler wear track which results in the increasing coefficient of friction. A much more minor

upward trend in the coefficient of friction for graphite observed under the same conditions.

Because of the number of wear tests completed, the findings of the graphite- $\mathrm{Sb}_2\mathrm{O}_3$ study are not conclusive, however, support is provided for the additive deformation concept.

CHAPTER 1X DISCUSSION

The role of bulk additions to solid lubricants may be one of simply permitting the formation of an improved tribological surface by deformation of the additive at or near asperity contact which results in densification or sintering of the surface. Since tribological surfaces are complex, it is difficult to provide direct evidence of such a role; however, qualitative improvement of compacts which contain beneficial additions are easily observed either by optical or scanning electron microscopy. This effort shows that the several alternative hypotheses proposed to date cannot be supported, and that the hypothesis advanced is consistent with observations in this effort and with those of others (8,10) in studies of the MoS2-Sb2O3 system.

Harmer and Pantano⁽⁸⁾ conclude that a role of $\mathrm{Sb}_2\mathrm{O}_3$ might be to retard oxidation of MoS_2 . However, oxidation studies of $\mathrm{Sb}_2\mathrm{O}_3$ here support those of others who have concluded that sublimation of $\mathrm{Sb}_2\mathrm{O}_3(c)$ occurs simultaneously with oxidation which rationalizes the thermal behavior of $\mathrm{MoS}_2\mathrm{-Sb}_2\mathrm{O}_3$ compacts without hypothesising that $\mathrm{Sb}_2\mathrm{O}_3$ in some way retards MoS_2

oxidation. Data do indicate that the oxidation rates of finally ground Sb203(c) or Sb203(c), or mixtures are related to equilibrium sublimation pressure in a pseudo-isobaric reaction.

DTA data also indicate that the exidation rate is dependent upon Sb203 surface area, which is as expected for an exidation rate dependent on sublimation of Sb $_{\Delta}$ 06.

Such a reaction is supported by calculation of free energies which indicate a greater driving force for oxidation of ${\rm Sb_40_6}$ than ${\rm Sb_20_3}(s)$.

Development of a more adequate understanding of the oxidation of ${\rm Sb}_2{\rm O}_3$ indicates that Harmer and Pantano's conclusions regarding a proposed protective role for ${\rm Sb}_2{\rm O}_3$ in retarding oxidation of ${\rm MoS}_2$ cannot be supported.

In DTA studies more directly related to lubricant compacts, when Sb_2O_3 was mixed with MoS_2 , no significant influence on oxidation of MoS_2 by Sb_2O_3 was found, suggesting that the role proposed for Sb_2O_3 to oxidize sacrificially, preventing oxidation of MoS_2 , cannot be supported. When MoS_2 was oxidized in DTA, it was found that the oxidative exotherm scarted at about 345°C (DTA 181), while the oxidation endotherm began at about 415°-420°C (DTA 175). Thus, MoS_2 begins to oxidize before Sb_2O_3 and, therefore, cannot be protected sacrificially by Sb_2O_3 for love mixtures of powders as avaluated in DTA.

However, Sb_2O_3 may promote densification of the

tribological surface. Dense packing of $8b_20_3$ around HoS_2 could retard the diffusion of oxygen into subsurface HoS_2 sites and would certainly be beneficial.

One might propose that addition of Sb2O3 to MoS2 allows the compact to reach closer to theoretica density at relatively low loads. Compaction of MoS2-Sb202 (<30 µM) and $MoS_2 = MoO_3$ (<30 μ M) formulations at modest pressures (=172 MPa) indicate that additives do not aid in achieving theoretical density. In fact, a smaller fraction of theoretical density was achieved for MoS2 with additives than for MoS_2 alone. The data are in contrast to Harmer and Pantano's (8) results in which there was modest improvement in compaction on a theoretical basis when Sb₂O₃ was added to MoS2. Data from this work show that the fraction of theoretical density achieved linearly decreased with increasing additive concentrations. The differences are not inconsistent if the Sb2O3 used by Harmer and Pantano contained large masses with voids which collapsed under high loading, or if their MoS2 was particularly difficult to densify.

The data generated in this effort suggest that densification of MoS_2 is not aided by bulk additions and a relatively low pressure. If, indeed, surface densification of MoS_2 films containing selected oxides and sulfides does occur during contact, as proposed then tribological conditions must generate asperity contact temperatures

sufficient to deform the additive, allowing effective densification of the surface layer.

It has been suggested that a sutectic of \$b203-MoO3 forms at tribological interfaces. (7) A eutectic compositon of these exides was prepared in the laboratory, and the eutectic endotherm was detected by DTA under argon. Interfacial materials removed from the sliding interface of compacts and debris gathered in the area of sliding contact generated at room temperature and at 370°C during wear tests were analyzed by DTA and x-ray diffraction techniques. No evidence of Sb₂O₃-MoO₃ eutectic formation was found. After sliding at room temperature, compacts showed a slight broadening of the most intense MoS2 reflection, and Sb₂O₃ peaks were lower in intensity, This may be due to MoS2 coverage of much of the interface after sliding. After wear at 370°C, x-ray spectra indicated that the dimorph $5b_20_3(c)$, as added, changed to $5b_20_3(o)$, and suggested trace oxidation of $\mathrm{Sb}_2\mathrm{O}_3$ to $\mathrm{Sb}_2\mathrm{O}_4$. Therefore, the concept that improved tribological properties are the result of eutectic formation cannot be supported by these analyses. Additionally, previous we set data (7) in air and vacuum and data of this work for 232°C wear tests indicate that MoS2 containing Sb203 is tribologically superior to MoS₂ alone whether evaluated in air, vacuum or inert atmosphere. These data do not support of a role in which

exidation of either ${\rm Sb}_2{\rm O}_3$ or ${\rm HoS}_2$ is required for the additive effects of ${\rm Sb}_2{\rm O}_3$ additions to be beneficial.

In other wear studies, it was found that additions of MoO₃ are superior to 8b₂O₃ additions to MoS₂ compacts when worn at 316°C in air. These data prove that the benficial role of Sb₂O₃ as an additive to MoS₂ is not unique.

Additionally, mixtures of MoO₃-Sb₂O₃ as MoS₂ additions are linearly intermediate in tribological benefit. Therefore, there does not appear to be beneficial synergism of MoO₃-Sb₂O₃ additions. Such synergism should be expected if formation of an MoO₃-Sb₂O₃ eutectic performs a significant tribological role in MoS₂-Sb₂O₃ solid lubricant compacts.

When other oxides and sulfides are added to ${\rm MoS}_2$ on a volumetric basis equivalent to 25 mole percent ${\rm Sb}_2{\rm O}_3$ addition, many of the compacts were found to be tribologically superior to pure ${\rm MoS}_2$. Additives that resulted in improved tribological performance appeared to have smoother compact surfaces when examined by both optical and scanning electron microscopy. This improvement in tribological condition could not be quantified but was easily observed. For the additions investigated, there is an excellent correlation of wear volume with additive shear strength. These data also support a concept that ${\rm Sb}_2{\rm O}_3$ is not unique in its ability to improve the tribological performance of ${\rm MoS}_2$.

For the orides and sulfides used in this study as compact additions, a transition from brittle to tough behavior should occur at 0.5 to 0.7 of the absolute melting temperatures (98) of those materials. Comparison of computed transition temperatures for the additions to MoS₂ suggests that such a temperature region of ductility is schievable for the beneficial additives. Based upon calculations for the MoS₂-Sb₂O₃ lubricant compact system, deformation of Sb₂O₃ at asperity contact is certainly possible, since temperatures in the region and above 0.5 to 0.7 of Sb₂O₃ melting temperature are achieved.

An evaluation of the wear of graphite- $\mathrm{Sb}_2\mathrm{O}_3$ compacts in which additions of $\mathrm{Sb}_2\mathrm{O}_3$ marginally improve graphite performance at temperatures lower than $\mathrm{MoS}_2\mathrm{-Sb}_2\mathrm{O}_3$. The properties of graphite and MoS_2 affect asperity contact temperatures resulting in higher asperity contact temperatures for the graphice- $\mathrm{Sb}_2\mathrm{O}_3$ system compared with $\mathrm{MoS}_2\mathrm{-Sb}_2\mathrm{O}_3$. Therefore, the beneficial tribological effect is found at a lower test temperature for graphite- $\mathrm{Sb}_2\mathrm{O}_3$. These data support a hypothesis that MoS_2 is not a unique solid lubricant in its role in $\mathrm{MoS}_2\mathrm{-Sb}_2\mathrm{O}_3$ formulations.

Thus, it appears that the beneficial mechanism of ${\rm Sb_2O_3}$ addition to ${\rm MoS_2}$ is not unique, neither for additive or solid lubricant. It does appear that the shear strength of the additive is critical in performance of compacts under tribological conditions, which supports the concept

position for the solid lubricant resulting in overall improved performance.

Excellent correlation of additive shear strength with wear volume and poorer correlation with friction coefficient supports the hypothesis in that critical beneficial deformation for solid lubricant accommodation occurs in the bulk of the compact and not at the interface.

CHAPTER X

CONCLUSIONS

Three main hypotheses have been proposed to date to rationalize the beneficial effects of ${\rm Sb}_2{\rm O}_3$ additions to ${\rm HoS}_2$. They are:

- (1) Sb203 oxidized secrificially, rather than MoS2.
- (ii) Sb_2O_3 retards the oxidation of MoS_2 by some ill-defined protective mechanism.
- (iii) A sutectic of $8b_2o_3$ -MoO₃ forms, aiding transfer film formation.

These hypotheses cannot be supported based upon the following Sheervations:

- (1) Initial exidation of MoS2 occurs at lower temperatures than for Sb2O3.
- (ii) Sublimation of Sb_40_6 appears to be a critical step in the oxidation mechanism of Sb_20_3 , which aids in rationalizing observations of others regarding the effects of Sb_20_3 on MoS_2 oxidation.

- (111) Bulk additions of $8b_2c_3$ to $Ho8_2$ are beneficial whether such compacts are evaluated in vacuum, inert atmosphere or air.
- (14) DTA or x-ray diffraction evidence of 8b203-NoO3 entectic formation or wear surfaces could not be found, although such a entectic could be prepared and detected in the laboratory.
- (v) Mixtures of ${\rm MoO_3}$ and ${\rm Sb_2O_3}$ as bulk additives to ${\rm MoS_2}$ did not act synergistically to provide tribological benefits greater than either of the additives individually.

Alternatively, it is proposed that the beneficial effect of $\mathrm{Sb}_2\mathrm{O}_3$ additions to MoS_2 is the result of deformation of $\mathrm{Sb}_2\mathrm{O}_3$ at asperity contact temperatures to permit and retain a tribologically preferred orientation of the solid lubricant. This hypothesis is supported by the following observations:

- (1) Bulk, rather than trace quantities of the additive are required.
- (11) SEM micrographs show that $MoS_2-Sb_2O_3$ compact wear surfaces appear to be much smoother and more dense compared with MoS_2 compact wear surfaces.

- -(111) MoO3 as an additive for MoS2 is superior to Sb2O3 with regard to wear volume and is equivalent in comparison of friction coefficient.
 - (1v) Other stable oxides and sulfides are also beneficial as additives to MoS2 compacts when evaluated up to 316°C in air; thus, Sb203 is not a unique additive.
 - (v) Limited wear studies of graphite-Sb₂O₃ compacts suggest that the beneficial effects of additives are not unique to MoS₂.
 - (vi) The low wear volumes of oxide and sulfide additions to MoS₂ appear to be related strongly to additive shear strength at test temperature. Correlation of wear volume with basic properties of the additive such as Moha hardness or melking temperature is not as good. The weaker correlation is probably due to the limited utility of using such properties as predictors of shear strength.
- (vii) Basic properties of the additives could not be used to predict coefficients of friction of compacts containing additives. The poor correlation of frictional coefficient with basic additive properties such as shear strength, Hohs hardness and melt temperature

compared to excellent to good correlation of those properties with wear volume indicates that the additions are not beneficial at the interface rather in the bulk.

This conclusion supports the hypothesis that the additives deform to permit and retain a tribologically preferred orientation for the solid lubricant.

These observations suggest that a beneficial additive must be thermally and chemically stable at such temperatures, yet shear to permit deformation at asperity contact temperatures. Thus, although many additives may be beneficial, selection of the solid lubricant, bulk additive and tribological conditions must be considered to define an optimum compact system.

REFERENCES

- N. N. Gardos and B. D. McConnell, "Development of a High-Load, High-Temperature Self-Lubricating Composite -- Part III: Additive Selection," ASLE-ASME Lubrication Conference, October 5-7, 1981, New Orleans, LA.
- 2. J. W. R. Taylor, ed., Jane's All the World's
 Aircraft, Jane's Publ. Co., New York, NY, 1984, p.
 893.
- 3. G. P. Hurphy, Jr., <u>U.S. Patent 3,314,885</u>, April 18, 1967 (CA 67: 1265t).
- 4. M. Campbell and V. Hopkins, "Development of Polyimide Bonded Solid Lubricants," <u>Lub. Eng.</u>, 23, 7 (1967).
- 5. R. D. Hubbell and B. D. McConnell, "Wear Behavior of Polybenzinidezole Bonded Solid-Film Lubricants," J. Lub. Tech., 92, Series F, 2 (1970).
- 6. B. D. McConnell, "Solid Lubricating Films Friction-Wear Improvement with Poly(benzothiazole) Binders," Ind. Eng. Chem. Prod. Res. Develop., 9, 4 (1970).
- 7. M. T. Levik, R. D. Hubbell and B. D. McConnell, "Oxide Interaction A Concept for Improved Performance with Molybdenum Disulfide," <u>Lub. Eng.</u>, 31, 20-27 (1975).
- 8. R. S. Harmer and C. G. Pantano, "Synergistic Mechanisms of Solid Lubricants," AFML-TR-77-227, December 1978.
- 9. M. I. Nosov, "The Lubricating Properties of Molybdenum Disulfide with Antimony Oxides and Sulfides," Khim. Tech. Topliv Mosel, 7, 43-44 (1978).
- 10. E. Yamamoto, K. Wada, T. Fukuzuka, K. Shimogori, K. Fujiwara and K. Tanji, "Lubricating Films to Prevent Galling of Stainless Steel Threaded Parts," <u>Lub. Eng.</u>, 40, 588-597 (1984).

11. F. J. Clauss, Solid Lubricants and Self-Lubricating Solids, Academic Press, New York, NY, 1972, pp. 1-4.

- 12. F. P. Bowden and D. Tabor, The Friction and Lubrication of Solids, Part II, Oxford Univ. Press, 1964.
- 13. R. Rabinowicz, Friction Wear of Materials, Wiley, New York, NY, 1965, pp. 114-117.
- 14. J. F. Archard, "Contact and Rubbing of Flat Surfaces," J. Appl. Phys., 24, 981-988 (1953).
- 15. A. Cameron, The Principles of Lubrication, John Wiley and Sons, New York, NY (1966).
- W. E. Campbell, "Solid Lubricants," in Boundary Lubrication, An Appraisal of World Literature, F. F. Ling, E. E. klaus, and R. S. Fein, Pub. by ASME, New York, NY, 1969.
- 17. W. O. Winer, "Molybdenum Disulfide as a Lubricant: A Review of the Fundamental Knowledge," Wear, 10, 422-452 (1967).
- 18. P. M. Magie, "A Review of the Potentials of the New Heavy Metal Derivative Solid Lubricants," Lub. Eng., 22, 262-269 (1966).
- 19. W. E. Jamison, "Structure and Bonding Effects on the Lubricating Properties of Crystalline Solids," ASLE Trans., 15, 296-305 (1972).
- 20. W. E. Jamison, Final Report on Solid Lubricant Research to Hughes Aircraft, Contract F33615-78-C-5196, October 1980.
- 21. A. Madhukar, "Structural Classification of Layered Dichalcogenides of Group IVB, VB and VIB Transition Metals," Solid State Comm., 16, 383-388 (1975).
- 22. W. E. Jamison and S. L. Cosgrove, "Frictional Characteristics of Transition Metal Disulfides and Diselenides," ASLE Trans., 10, 19-27 (1967).
- 23. D. J. Boes, "Unique Solid Lubricating Materials for High Temperature Applications," ASLE Trans., 10, 19-27 (1967).

- 24. D. J. Boes and B Chamberlain, "Chemical Interactions Involved in the Formation of Oxidation-Resistant Solid Lubricant Composites," ASLE Trans., 11, 131-139 (1968).
- 25. R. D. Dayton, "Experimental Investigation of Ag-Hg-W7FE-MoSe₂ Solid Lubricated Ball Bearings for High Speed, High Temperature, and High Load Applications," <u>AFAPL-TR-71-100</u>, January 1972.
- 26. D. H. Killeffer and A. Ling, Molybdenum Compounds:
 Chemistry and Technology, Interscience, New York, NY,
 1952.
- 27. C. A. Hampel, Uncommon Metals, Reinhold Publ. Co., 283 (1961).
- 28. R. C. Weast, ed., <u>Handbook of Chemistry and Physics</u>, 61st ed., Chemical Rubber Co., Cleveland, OH, 1981.
- 29. M. M. Spivak, A. S. Pashinkin, R. A. Isakova and K. S. Amosova, "Thermodynamic Analysis of the Mo-S-O System," Zhin. Prik. Khin, 51, 549-553 (1978).
- 30. R. G. Dickinson and L. Pauling, <u>J. Am. Chem. Soc.</u>, 45, 1466 (1923).
- 31. C. R. Cardoen, "The Kinetics of Molybdenum Disulfide Oxidation," Doctoral Disseratation, Univ. of Utah, 1969.
- 32. N. B. Pilling and R. G. Bedworth, <u>J. Inst. Hetals</u>, <u>29</u>, 532 (1922).
- 33. A. N. Zelikman and L. B. Belaevskaya, "Concerning the Melting Temperature of Molybdenite," J. Inorg. Chem (USSR), 10, 2245-2256 (1956).
- D. H. Buckley, "Examination of Molybdenum Disulfide with LEED and Auger Emission Spectroscopy," NASA TN D-7010, December 1970.
- 35. W. E. Swartz and D. M. Hercules, "X-Ray Photoelectron Spectroscopy of Molybdenum Compounds," Anal. Chem., 43, 1774 (1971).
- 36. L. B. Atkinson and P. Swift, "A Study of the Tribo-Chemical Oxidation of Molybdenum Disulfide Using X-Ray Photo-Electron Spectroscopy," Wear, 29, 129-133 (1974).

- 37. F. Lepage, P. Baillif and J. Bardolle, "Study of Electron Spectrometry (ESCA) of the Oxidation States of Molybdenum," C. R Hcbd. Seances Acad. Sci., Series C, 280: 1089-1092 (1975).
- 38. F. Lepage, P. Baillif and J. Bardolle, "Study of Various Degrees of Oxidation of Molybdenum and Tungsten by Electron Spectroscopy (ESCA)," Vide, Series: 30A, 2nd Colloq. Int. Phys. Chim. Surfaces, 101-103 (1975).
- 39. S. O. Grim and Luis J. Matienzo, "X-Ray Photoelectron Spectroscopy of Inorganic and Organometallic Compounds of Molybdenum," <u>Inorg. Chem.</u>, <u>14</u>, 1014-1018 (1975).
- 40. J. M. Wilson, "Energy Loss Spectroscopy of Sulfide Films on Mo(100)," Surf. Sci., 57, 499-508, (1976).
- 41. T. I. Lin and D. Lichtman, "AES Studies of Chemical Shift and Beam Effect on Molybdenum Oxides," J. Vac. Sci. Tech., 15, 1689-1694 (1978).
- 42. Y. Toga, A. Isogai, and K. Nakajima, "LEED-AES Study of Sulfide Single Crystals," <u>Surf. Sci.</u>, <u>86</u>, 591-600 (1979).
- C. Y. Wang, Antimony: Its Geology, Metallurgy,
 Industrial Uses and Economics, C. Griffen and Co.,
 London, 1952.
- 44. D. J. Stewart, O. Knop, C. Ayasse and F. W. D. Woodhams, "Pyrochlores VII. The Oxides of Antimony: An X-Ray and Mossbauer Study," Can. J. Chem., 50, 690-700, (1972).
- 45. H. Remy, Treatise on Inorganic Chemistry, Vol. 1, Elsevier, New York, NY, 1956.
- 46. E. J. Roberts and F. Fenwick, "The Antimony-Antimony Trioxide Electrode and Its Uses as a Measure of Acidity," J. Am. Chem. Soc., 50, 2125-2147 (1988).
- W. B. White, F. Dachille and R. Ray, "High-Pressure Polymorphism of As₂O₃ and Sb₂O₃," Z. Krist., 125, 450-458 (1967).
- 48. P. Gopalakrishnan and H. Manohar, "Kinetics and Mechanism of the Transformation in Antimony Trioxide from Orthorhombic Valentinite to Cubic Senarmontize,"

 J. Solid State Chem., 15, 61-67 (1975).

- 49. W. B. Hincke, "The Vapor Pressure of Autimony Trioxide," J. Amer. Chem. Soc., 3869-3877 (1930).
- 50. E. Jungermann and K. Plieth, "Dampfdrucke und Kondensations-geschwindig-keiten der polymorph Arsen-und Antimontiroxide," Z. Phys. Chem. Neue Folge, 53, 215-228 (1967).
- 51. C. G. Naier, U.S. Bur. Mines Rep. Investigations, Nr. 3262, December 1934, pp. 22-28.
- 52. C. A. Cody, L. DiCarlo and R. K. Darlington,
 "Vibrational and Thermal Study of Antimony Oxides,"

 <u>Inorg. Chem.</u>, <u>18</u>, 1572-1576 (1979).
- 7. K. Agrawal, A. L. Shashimohan and A. B. Biswas, "Studies on Antimony Oxides: Part 1, Thermal Analysis of Sb₂O₃ in Air, Nitrogen and Argon," J. Thermal Anal., 7, 635-641 (1975).
- P. Gopalakrishnan and H. Manohar, "Topotoxy in the Oxidation of Valentinite, Sb₂O₃, to Cervantite, Sb₂O₄," <u>Pramana</u>, 3, 277-285 (1974).
- 55. P. Gopalakrishnan and H. Manohar, "Topotactic Oxidation of Valentininte Sb₂O₃, to Cervantite, Sb₂O₄: Kinetics and Mechanisms," <u>Journal of Solid State Chemistry</u>, 16, 301-306 (1976).
- 56. S. E. Golunski, T. G. Nevell and M. I. Pope, "Thermal Stability and Phase Transitions of the Oxides of Antimony," Thermochimica Acta, 51, 153-168 (1981).
- 57. D. Rogers and A. C. Skapski, "The Crystal Structure of β-Sb₂O₄: A New Polymorph₆" Proc. Chem. Soc., 400 (1964).
- 58. T. Birchall, J. A. Conner and I. H. Hiller, "High-Energy Photoelectron Spectroscopy of Some Antimony Compounds," J. Chem. Soc., Dalton Trans., 20, 2003-2006 (1975).
- 59. F. Garbassi, "XPS and AES Study of Antimony Oxides,"
 Surf. and Interface Anal., 2, 165-169 (1980).
- 60. C. D. Wagner, "Chemical Shifts of Auger Lines, and the Auger Parameter," Faraday Discuss., 60, 291-300 (1975).

- 61. V. Hopkins and M. Campbell, "Friction and Wear-Life of Selected Solid Lubricant Films at -100°F, R.T., and 400°F," Lub. Eng., 430-435 (1969).
- 62. A. E. Standage and W. N. Turner, "A High-Temperature Polyimide Reinforced with Silica Fiber," J. Materials Sci., 2, 103-111 (1967).
- R. L. Fusaro, "Polyimides-Tribological Properties and Their Use as Lubricants," in K. L. Mittal, ed.

 Polyimides: Synthesis, Characterization and Applications, Vol. II, Plenum Press, New York, NY, 1982. p. 1066.
- 64. M. Parmentier, A. Courtoir and C. Gleitzer, Bull. Soc. Chem. Fr., 1-2 (Pt 1) 75-77 (1974).
- 65. J. B. Peace, "Solid Lubricants" in E. R. Braithwaite, ed., <u>Lubrication and Lubricants</u>, Elsevier, Amsterdam, 1967, p. 95.
- 56. J. B. Peace, "Solid Lubricants" in E. R. Braithwaite, ed. <u>Lubrication and Lubricants</u>, Elsevier, Amsterdam, 1967, p. 72.
- J. W. Lyons, The Chemistry and Uses of Fire Retardants, Wiley-Interscience, New York, NY, 1970, pp. 209-218.
- 68. D. A. Church and F. W. Moore, "Molybdenum-Based Flame Retardants are Now in the Plastics Ball Game," Plast.

 <u>Eng.</u>, 31, 36 (1975).
- 69. D. Edelson, V. J. Kuck, R. M. Lum, E. Scalco, W. H. Starnes and S. Kaufman, Jr., "Anomalous Behavior of Molybdenum Oxide as a Fire Retardant for Polyvinyl Chloride," Comb. and Flame, 38, 271 (1980).
- 70. O. Borger and J. Krogh-Moe, "The Infrared Spectra of Some Modifications of Arsenic Trioxide and Antimony Trioxide, Acta Chem. Scand., 11, p. 265-267.
- 71. J. R. Ferraro, J. S. Ziomek and G. Mack, Spectrochem.
 Acta, 17, 802 (1961).
- 72. V. G. Trofimov, A. I. Sheinkman and G. V. Kleshehev, "Griented Crystallization of Cervantite on Heating Sb₂O₃," <u>Izvestiga VSZ</u>, <u>Fizika</u>, 3, 135-137 (1973).

73. I. Langmuir, Physik, Z., 14, 1273 (1913).

- 74. P. J. Durrant and B. Durrant, <u>Introduction to Advanced Inorganic Chemistry</u>, Wiley, NY, 1970, p, 773.
- 75. J. B. Peace, "Solid Lubricants" in E. R. Braithwaite, ed. <u>Lubrication and Lubricants</u>, Elsevier, Amsterdam, 1967, p. 76.
- 76. M. E. Campbell, "Solid Lubricants, A Survey," NASA SP-5059(01) National Aero Space Adm., Washington, DC, 1972.
- 77. J. B. Peace, "Solid Lubricants" in E. R. Braithwaite, ed. in <u>Lubrications and Lubricants</u>, Elsevier, Amsterdam, 1967, p. 91.
- 78. P. G. Lenhert, J. F. O'Brien, W. W. Adams, "A User's Guide to the Picker Diffractometer for Polymer Morphology Studies," AFWAL-TR-86-4024, Air Force Materials Laboratory, Wright-Patterson Air Force Base, OH, July 1985.
- 79. D. P. Anderson, "X-ray Analysis Software: Operation and Theory Involved in Program DIFF," AFWAL-TR-85-4079, Air Force Materials Laboratory, Wright-Patterson Air Force Base, OH, June 1985.
- 80. W. L. Bragg, "The Crystalline State," MacMillan, New York, 1933.
- 81. "Alphabetical and Grouped Numerical Index of X-ray Diffraction Data," American Society of Testing Materials, Special Technical Publication, No. 48E (1955).
- 82. G. Salomon, A. W. J. DeGee, and J. H. Zaat, "Mechano-Chemical Factors in MoS₂ Film Lubrication," in Mechanisms of Solid Friction, P. J. Bryant, M. Lavik and G. Salomon (Eds.), Elsevier, Amsterdam, 1964, p. 87.
- 83. A. DiSapio, "Bonded Coatings Lubricate Metal Parts,"
 Product Engineering, 31, 48-53 (1960).
- 84. M. B. Peterson and R. L. Johnson, "Friction Studies of Graphite and Mixtures of Graphite with Several Metallic Oxides and Salts at Temperatures to 1000°F," NACA TN3657, Dec 1955.
- 85. J. M. Palacios, A. Rincon and L. Arizmendi, "Extreme Pressure Lubricating Properties of Inorganic Oxides," Wear, 60, 393-399 (1980).

- American Standard for Testing Material Test Method D2783-82, "Neasurement of Extreme Pressure Properties of Lubricating Fluids (Four-Ball Method)," Vol. 05.02, Petroleum Products and Lubricants, ASTM Philadelphia, PA, 1983.
- B. Tabor, "Mohs's Hardness Scale A Physical Interpretation," Phys. Soc. Proceed. (London), B-67, 249-257 (1954).
- 88. E. Rabinowicz, <u>Friction</u>, <u>Wear and Lubrication</u>:
 <u>Tribology I</u>, MIT Press, Cambridge, MA, 1981, p. 1.12.
- 89. M. B. Peterson, S. B. Calabrese and B. Stupp, "Lubrication with Naturelly Occurring Double Oxide Films," DARPA Report, AD 124248 (1982).
- 90. T. F. J. Quinn and W. O. Winer, "The Thermal Aspects of Oxidational Wear," Wear, 102, 67-80 (1985).
- 91. M. B. Peterson, S. F. Murray and J. J. Florek, "Consideration of Lubricants for Temperatures Above 1000°F," <u>ASLE Transactions</u>, 2, pp. 225-234 (1960).
- 92. M. B. Peterson, "High Temperature Lubrication,"

 Proceedings of Lubrication and Wear, D. Muster and B. Sternlicht, eds., McCutchan Publishing Co., Berkely, CA, 1965.
- 93. E. Rabinowicz, <u>Friction Wear of Materials</u>, Wiley, New York, NY, 1965, pp. 87-89.
- 94. J. F. Archard. "The Temperature of Rubbing Surfaces," Wear, 2, 438-455 (1958/59).
- P5. R. H. Perry and C. H. Chilton, eds., Chemical Engineers Handbook, 5th Ed., McGraw-Hill, New York, NY, 1973.
- 96. H. Ernst and N. E. Merchant, "Surface Friction of Clean Metals A Basic Factor in the Metal Cutting Process," Proceedings of the Special Summer Conference or Friction and Surface Finish, Mass. Inst. of Tecn., Cambridge, MA, pp. 76-101, 1940.
- 97. C. Palache, H. Berman and C. Fronkl, "The System of Mineralogy of James Dwight Dana and Edward Salisbury Dana, Yrle University 1837-1892," 7th ed., Wiley, New York, AY, 1955.
- 98. C. E. Dieter, Mechanical Metallurgy, McGraw-Hill, New York, NY, 1976, p. 496.

- 99. M. B. Peterson, J. J. Florek and R. E. Lee, "Sliding Characteristics of Metals at High Temperatures," ASLE Transactions, 3, 101-109 (1960).
- 100. E. R. Braithwaite, Solid Lubricants and Surfaces, McMillan, New York, NY, 1964, p. 159.

- 101. P. Fleischauer and R. Bauer, "The Influence of Surface Chemistry on MoS₂ Transfer Film Formation," ASLE Preprint No. 86-AM-5G-2, Amer. Soc. of Lub. Eng. Annual Meeting, Toronto, Ont., May 1986.
- 102. J. B. Peace, "Solid Lubricants" in E. R. Braithwaite, ed., <u>Lubrication and Lubricants</u>, Elsevier, Amsterdam, 1967, p. 90.

APPENDIX A

PROCEDURE FOR ISOTHERMAL OXIDATION OF Sb203

- 1. The initial rig conditions are as follows:
 - a) All valves closed. Record atmospheric pressure and temperature.
 - b) Furnace lowered.
 - c) P2O5 tube installed in O2 supply line.
- 2. Weigh Sb202 and record weight (0.1 milligram).
- 3. Using stopcock fitting, mount Sb203 sample in rig.
- 4. Bring Hg level to that of the gas buret valve by opening system went and Hg bulb manipulation. Close buret valve bring system to atmospheric pressure.
- 5. Slowly open vacuum valve and evacuate system to ~1 torr. Close vacuum valve, and by opening 02 valve bring system to atmospheric pressure.
- 6. Repeat Step 5 two additional times.
- 7. Fill a gas buret wih 0_2^{+} to atmospheric pressure using the 0_2 valve. Loosen gas valve.
- 8. Use vacuum valve to reduce system pressure to 140 torr, typically, or other pressure desired.
- 99.99 percent 0_2 , research grade, Matheson, East Rutherford, NJ introduced through P_2O_5 drying tube.

- 9. Bring furnace into test position and start heating. When test some temperature has reached equilibrium at the desired temperature, again bring 02 pressure to 140 torr then drop sample into reaction some. Start clock and record sample temperature.
- 10. As exidation proceeds, 0₂ pressure in the system is reduced. Periodically, using the gas buret valve, gently bring the system 0₂ pressure back to 140 torr. Record the volume of 0₂ remaining in the buret at atmospheric pressure by Hg bulb manipulation.

 Record 0₂ vol/time data until calculated volume of 0₂ is consumed or for a period of up to two (2) hours.

 After 0.5 hour, if no more than 8 cm³ of 0₂ has been consumed, take reading volumetric at five or ten minute intervals.
- 11. At test termination, turn off and lover furnace.
- 12. When reaction tube is cool, bring system to atmospheric pressure with air. Remove and label sample. Use a new tube for the next determination.

APPENDIX B

ISOTHERMAL Sb203 OXIDATION DATA

		OXYGEN CONSUMPTION	
	TIME (SEC)	HG O2 & STP/g Sb2O3	√TIME (SEC ^{1/2})
Run #52	1441	4.39	38.0
\$b203(c)		5.38	40.74
30203(5)	2023	10.09	45.0
521°C	2175	12.23	46.64
321 0	2225	12-67	47.20
	2378	15.30	48.8
	2501	17.49	50.01
	2754	20.56	52.5
	2869	22.76	53.60
	3072	26.70	55.41
	3110	27.14	55.81
	3297	29.02	57.4
	3447	31.96	58.7
	3520	32.40	59.3
	3686	34.11	60.7
	4000	37.62	63.3
Run #55	515	3.456	22.7
\$b203(c		5.529	25.1
00203	779	8.985	27.9
542°C	869	11.749	29.5
	940	14.514	30.7
	993	15.896	31.5
	1056	18.660	32.5
	1100	21.424	33.2
	1147	23.498	33.9
	1189	25.571	34.5
	1225	28.336	35.0
	1273	30.409	35.7
	1325	33.174	36.4
	1402	37.200	37.4
	1475	40.085	38.4
	1539	42.158	39.2
	1628	44.923	40.4
	1698	46.996	41.2
	1815	48.378	42.6
	1928	49.760	43.9
	2218	51.834	47.1

mg 0_2 0 STP/g $8b_20_3$ /TIME (SEC^{1/2}) TIME (SEC) RUN #57 333 2.469 8b203(c) 641 6.172 698 9.875 542°C 745 13.578 COATED TUBE 812 19.750 879 24.069 935 26.538 1022 29.007 RUN #58 456 6.347 21.6 8b203(c) 527 9.520 23.0 14.28 25.3 641 559°C 731 22.214 27.0 800 28.3 26.974 862 31.734 29.4 941 36.494 30.7 1036 42.841 32.2 1138 33.7 46.015 1260 35.5 49.187 **RUN #59** 341 6.098 18.5 461 11.518 21.5 Sb203 527 15.583 23.0 579°C 587 21.004 24.2 648 25.5 27.778 687 30.489 26.2 743 35.910 27.3 802 41.329 28.3 916 30.3 49.461 32.1 1031 53.526 RUN #60 36.9 1364 5.395 Sb203(0) 1802 8.631 42.5 2240 47.3 11.328 453°C

12.408

16.184

18.341

49.4 57.1

67.8

OXYGEN CONSUMPTION

2442

3260

OXYGEN CONSUMPTION
TIME (SEC) mg 02 @ STP/g Sb203 /TIME SEC1/2

RUN #61	467	6.886	21.6
8b203(o)	627	10.329	25.0
	780	14.92	27.9
506°C	913	18.364	30.2
	1053	21.233	32.4
	7580	24.102	34.4
	1301	26.715	36.1
	1421	29.267	37.7
	1518	31.562	39.0
	1661	34.432	40.8
	2155	39.597	46.4
	2800	43.61	52.9
RUN #62	691	4.721	26.3
8b203(o)	836	7.419	28.9
	941	9.441	30.7
492°C	1124	12.814	33.5
	1304	15.848	36.1
	1568	20.232	39.6
	1827	24.953	42.7
	2151	29.336	46.4
	2551	34.392	50.5
	2690	36.417	51.9
	3780	41.475	61.5
	4672	43.161	63.4
	6000	44.510	77.5
RUN #63	308	9.587	
Sb, 03(0)	387	12.143	
2 3 1 2	478	17.256	
521°C	517	28.534	
	710	26.204	
	830	31.317	
	1060	38.347	
	1489	43.780	
	3129	46.656	

OXYGEN CONSUMPTION TIME (SEC) $mg O_2 Q STP/g Sb_2O_3 /TIME (SEC^{1/2})$

RUN #64	274	6.386	16.6
2 b ₂ O ₃ (o)	373	11.298	19.3
	438	14.245	20.9
521°C	524	18.667	22.9
	600	22.597	24.5
	735	28.000	27.1
	847	31.685	29.1
	1006	37.334	31.7
	1156	40.772	34.0
	1645	44.702	40.6
	2375	46.422	48.7
RUN #68	1495	6.405	38.7
50%	2125	10.551	46.1
Sb203(0)	3145	15.449	56.1
50%	4467	21.854	66.8
sb₂0₃(c)	6168	30.144	78.5
492°C			
RUN #69	708	6.714	26.6
502	1130	13.092	33.6
Sb ₂ O ₃ (o)	1594	20.141	39.9
2-3	1878	24.841	43.3
50%	2390	32.555	48.9
Sb ₂ O ₃ (c)	3111	37.597	55.8
2 3	6200	41.626	78.7
520°C			
RUN #83	2315	16.545	48.1
50%	3196	21.715	56.5
Sb ₂ O ₃ (o)	3815	25.335	61.8
2-3	4640	28.437	68.1
50% Sb ₂ 0 ₃ (c 504°G		200.00	3332
RUN #84	887	11.75	29.8
50%	1036	14.20	32.2
Sb ₂ 0 ₃ (o)	1242	17.14	35.2
% - 3 · · ·	1475	20.56	38.4
50%	1690	23.99	41.1
8b ₂ 0 ₃ (c)	1967	27.91	44.4
£ 3'	2430	31.82	49.3
530°C	•		

APPENDIX C

DIFFERENTIAL THERMAL ANALYSIS SUMMARY HEATING RATE, 10°C/MIN.

	EAU LEAR		
	576		No peaks
	576		Gentle Endotherm
	808		
1:1	495, 407		
2:1	550		
2:1	516	652	
			No peaks
2:1	572		
	1	!	
Sb ₂ O ₃ (o) Ground 8 Min.	514	653 (8)	
203(o) Ground 16 Min.	808	656 (8)	
$\mathrm{Sb}_2\mathrm{O}_3(\mathrm{o})$ Ground 8 Min.	473	Diff	Diff. \$5203(0)
$\mathrm{Sh}_2\mathrm{O}_3(\mathrm{o})$ Ground 16 Min.	352, 464	Diff	Diff. \$5203(0)
Sb ₂ 04/Sb ₂ 03(c) 8 Min 3:1	485		
8 Min. 1:1	530		
$\mathrm{Sb}_2\mathrm{O}_4/\mathrm{Sb}_2\mathrm{O}_3(\mathrm{c})$ Ground 8 Min. 3:1	184		
₩ €		550 516 572 514 508 473 485 530 530	652 653 (8

APPENDIX C (CONT'D)

DIFFERENTIAL THERMAL ANANLYSIS SUMMARY

85 S 102 Sb ₂ 0 104 Sb ₂ 0 107 Sb ₂	$89^{13}(c) - 89^{03}(0)$				
	1	57:43			Each ground
			Shoulder 429	with	8 Min., Mixe
	Sb203(o), I Min Grind		493	651	
	Jerry Sec Girid		BOC	624	
	Sb203(0), 2 Kin Grind		475	;	
	FINE Sh203(0) Grind 8 Min		1	i	In argon,
					gentle Endother
131	Sb203(c)		584		
164	MoS ₂		451, 500(m)		
169	Sb203 Ground 16 Min.		558		
175	Sb ₂ 0 ₃ (Alfa)		456, 487(m), 548	8	
181	Mo S ₂		465, 521(m)		
182	75 Hole % Sb ₂ 0 ₃ (Alfa)				
	25 Mole % HoS ₂		479(m), 509, 566	999	
183	25 Hole X Sb ₂ 0 ₃ (Alfa)				
	75 Hole % Hos2		440, 496(m), 547	, 547	
237	50-50 WEZ MOS-Sb203(0)		446, 495(m)		
	Pellet (Sb203(o) Grind 8 Min)	nd 8 Min)			
238	Sh203(o) Pellet		475		
	(Sb203(o) Ground 8 Min)	•			
small, t	small, i = large, n = major	ALP!	** ALPA Sb ₂₀₃ Contains Both Uimorphs	Both Dime	rphs

APPENDIX D

	FEEDET	1.040	TEMPERATURE		TEST	COEF. OF	COEF. OF FRICESON	W.AR	•
TEST	Soff Control	(1)	د.	AFRUSPHERE	TIME	Ju Shc	IN MIR	VII, BME.	`.
-,	1	3	77	DRY ASK	4 HKS	ł	040.	1.175	
-	Mes. + Sea	=	77	DRY AIM	* EES	. O 18	140.	40. 504	
4	No.S Cer	3	7.7	SEV AND	S # # 9	# :·	0/0-	77770	
٠.	Has , + val (Crushed)	90	77	URY AIR	*48 *	740.	640.	187.0	
£	Hos, vval(Crushed)	5	11	BRY AIR	SHH 9	.032	. 040	H . 26 5	
~	Ho S.,	90	77	MAY AIR	4 1185	.056	. 655	0.175	
転	20 6.	=	23	BRY AIR	4 22.5	.059	. 1161	0. 116	
	Mus, eval (Crushed)	9	11	DRY AIR	4 BRS	.040	.045	0.250	
3	Ke-52	901	7.7	KET AIR	20 MIN	069.	:	AI.L	
=	Ecc. + See	-	73	UET AIR	× 11 ×	.155	:	11. 286	
~	Hos. See	901	7.7	VET AIR	S HIR	.245	;	0.74	
=	No.S.	=	2.2	UET AIR	- HIE	€ 1 ,	;	AL.	
7	Mo S 3	=	7.7	WET AIR	=======================================	177.	!	J. A.	
~	Kow, . val(Crashed)	ě	77	UET AIR	S MIR	9 8 7 .	;	9.1.0	
=	Nos, +Cer	3	22	WET AIR	> NIR	. 2015	;	0.92	
-	No.S.	00:	9	DRY AIR	- 11	91 0.	;	0.054	
<u>=</u>	, som	=	-	UET AIR	S R.I.R	.117	1	0.151	
2	Musz+Val(Cřushed)	<u>e</u>		DRY AIR	20 HIN	160.	;	06078	
=	Hus. +Cer	100	3	DRY ASE	20 MIN	071.	. 250	0.17	
~	8-84 Sul	25	541	DRY AIR	# I W 417	.150	.172	0/1-0	
-7	Hus, that	90	941	BET ASR	20 MIN	090.	.218	960.0	
*:	Hes.	001	160	VET AIR	ZU MIN	.125	.175	710.0	
\$	MoS,+Sen	201	04-	WET AIR	81H 07	. 106	. 151	7117	
47	Hes to Sen	100	. 160	WET AIR	ZU MIN	.085	751.	1.42	
ŧ	S C M	9	22	DRY AIR	20 1118	901.		0.1641	
Ξ	Mos.	001	22	DRY AIR	20 MIN	011.		0.1611	
=	Hos.+Kal	5	2.2	DRY AIR	20 MIN	390.	. 1167	6.052	
?	Husytan	100	22	DRY AIR	20 MIN	.055	.052	0.070	
=	Hns, ther	9	22	DRY AIR	20 MIN	740.	<u>:</u>	0.236	
7.1	Mo S. + C.P.	991	22	DRY AIR	ZO MIN	.050	697.	0.274	
5	Mos, sen	2	2.2	NET ASR	20 HIN			1.44.1	
Ę	(a) (b) (c)	50	23	WET AIR	MIN 07	=	8 1	107.0	
7	Suff.	=		Sit Rel. Hum.	# = T	6 60.	701.	11.176	
3	TRA+, SOM	-	212	Sot Rel. Hum.	H H	860.	. 11411	15.1141	
2	S of the second	100	717	DRY ASSOR	H H ==	. 120	160.	1:7:1	
3	Hos, eval	=	747		HIH OF	. 407.9	94,01	11. 11.3t	
,	Mox	===	717			. 44.7	580.	0.175	
. ,	Harry o Val	===	717	DRY ARCHN	N E	7 40.	.075	1:0:0	

	PELLET	LOAD	TEMPERATURE		TEST	COEF	OF	FRICTION	YEAR
TEST	# COMPOSITION	(g)	5 .	ATHOSPHERE	TIME	30 880	10 HIN	IS HIN	VOLUME (mm ³)
43	MoS,+Sen	100	232	DRY ARGON	NIM OI	.072	080.	1	2.206
44	MoS,+Cer	100		DRY ARGON	10 MIN	.079	001.	!	2.232
4.5	MoS ₂ +Sen	100	\mathbf{c}	DRY ARGON	10 MIN	.083	.083	!	90
9 %	MôS,	00 i	~	DRY ARGON	NIM OI	111.	_	!	0.352
47	Mos,	100	232	DRY AIR	0	.123	. 123	;	16
48	MoSy+Sen	100	~	DRY AIR	350 SEC	690	;	;	2.638
49	MosyrCer	100	3	DRY AIR	10 MIN	.080	.110	;	0.307
2 C	MoS2+Val	001	232	DRY AIR	NIM OI	.063	.063	1	0.024
51	Moss	100	~	WET AIR	10 MIN	. 110	.140	1	0.039
52	MoS	100	232	WET AIR	10 MIR	.130	.133		0.200
53	MoS2+Val	100		WET AIR	10 MIN	. 088	.075	1	0.126
54	MoS,+Cer	100	3	WET AIR	NIM OI	1	.133	1	0.265
55	MoS,+Sen	100	232	WET AIR	NIM OI	.022	080	i	1.780
96	MoS,+Sen	001	3	WET AIR	NIM OI	1900	.063	!	0.393
57	MoS,+Val	100	3	WET AIR	10 MIN	.087	.073	!	0.131
58	Hos,	100	3	DRY AIR	NIM OI	.018	.130	!	0.212
59	MoS,	100	232	DRY APGON	NIW OI	.104	.107	:	0.109
09	Mo S ₂	100	3	WET AIR	NIW OI	760.	.087	;	0.369
61	MoS,+Val	001	316	DRY AIR	NIM OF	.065	060.	1	0.193
62	MoS,	100	(474)	DRY AJR	10 MIN	.143	.150	1	0.364
63	Mos,+Sen	100	316	DRY AIR	10 MIN	.085	, 204		7.972
94	MoS,+Cer	160	316	DRY AIR	IO MIN	.077	.188	!	181.0
65	Mos	100	-	DRY ARGON	NIM OF	.108	100	1	0.641
99	MoS,+Val	100	_	DRY ARGON	10 MIN	.065	.085	;	0.125
19	MoS,+Cer	100	_	DRY ARCON	NIM OI	.075	.187	;	0.536
8 9	McS ₂ +Sen	001		DRY ARGON	NIW OF	.052	.137	:	5.652
69	MoS,+Cer(Crushed)	100	232	WET AIR	10 MIN	• 109	114	:	0.166
70	F MoS ₂	100	-	DRY AIR	IS MIN	.079	.142	.153	0.158

WEAR	VOLUME (DM3)	0.108	2.843	0.352	0.126	1.121	2 5	E	0,203	0.344	0.116	0.047	0.166	0.048	0.060	0.073	0.056	0.047	0.120	0.111	0.128	0.172	0.263	0.440	0.183	0.110		0.140
	15 MIN	0110	.116	!	!	;	264		890.	i	960 .	ł	.056															
	10 Min	100	.082	;	;	ł	.267	.259	.064	.029	90	ł																
	30 SEC	.062	.043	:	;	¦	.208		.073	.047	690.	.077	.056						•									
TEST	TIME	HIN	RIR	SEC	SEC	S C	SEC	SEC	SEC	HIN	SEC	SEC	SEC	SEC	SEC	SEC	SEC	SEC	SEC	SEC	SEC	SEC	SEC	SEC	SEC	SEC		SEC
	TI	18.3	22.8	1000	0001	000	1000		1000	01	1000	1000	1000	9	150	300	9	09	300	150	300	450	450	150	300	09		300
	ATHOSPHERE	AIR	-	DRY AIR		DRY AIR	AIR	AIR	AIR	DRY AIR		DRY AIR	DRY AIR	DRY AIR	DRY AIR	DRY AIR	VACUUM	DRY AIR	DRY AIR	DRY AIR	DRY AIR	DRY AIR	DRY AIR	DRY AIR	DRY AIR	DRY AIR		URY AIR
TEMPERATURE	¥ 0.			316	316	316						316	316	316		16	316	316	٠	16	16	316	316	316		316		316
LOAD	(g)	100	100	100	100	100	100	100	100	100	100	20	150	100	100	100	100	100	100	100	100	100	100	100	100	100		100
PELLET	COMPOSITION		MoS ₂ +Sen	0.5	HoS2/Val	Mos,/Sen	MoS,+Cer	85 MoS,/15 Val	65 MoS2/35 Val	Mos,/Sen/Val	MuS ₂ /Sen/Val	MoS2/Val	MoS ₂ /val	MoS2+Val	MoS2+Val	MoS2+Val	Mos ₂	Mos	Mos	MoS	MoS ₂	MoS ₂	MoS ₂ +Sen	MoS ₂ +Sen	%oS ₂ +Sen	MoS ₂ +Sen	MoS2+Val(Heated	Treated
	TEST	7.1	72	4 E	7.5	76	11	78	62	9.0		82	83	84	8 2	98	87	88	89	06	91	92	93	96	9.5	96	16	

ATB AGA			
274			0.167
			0.099
AIR 300			0.150
AIR 60			0.077
AIR 1000	•	.139	0.059
AIR 1000		690.	0.074
AIR 1000		.150	0.043
AIR 1000			0.087
AIR 1000			0.053
AIR 1000			0.037
AIR 1000			0.082
AIR 1000			0.079
AIR 1000		. 140	0.055
AIR 1000		001.	0.057
AIR 1000		. 113	0.075
AIR 1000		.086	0.181
AIR 1000		.118	0.134
ATR 1000		.115	0.105
AIR 1000		.171	0.045
AIR 1000		.129	0.057
AIR 1090		699.	0.031
AIR 1000		.154	•
AIR 1000		.158	0.050
AIR 1000		.136	0.081
DRY AIR 1000 SEC	. 052 . 083 . 072 . 072 . 075 . 075 . 109 . 1127 . 078 . 014		. 139 . 069 . 150 . 100 . 113 . 115 . 129 . 158 . 158

	PR1.1.87	LOAD	TEMPERATURE	٠		TEST	COEF. UF	FRICTION	VEAR
7267	7		(3.)	ATHOSPHERE	PHERE	TIME	30 SEC	15 MIN	VOLUNE (mm)
7	(a) 0 48+ 30M	1	316	DRY	AIR	1000 SEC	.078	.138	0.108
124	Mos. +54.0.(0)	_	316		AIR	•	.100	- 192	0.078
125	(0) -0 -4S+ -0 -0 + - 3 -0 M	-	316		AIR		.088	.178	0.079
136	Graphteatsh.O.(0)	-	316		AIR		901.	. 228	0.00
127	Graphic 523,57	-	316		AIR		.269	.304	0.043
1 2 0	Creshtretch. O. (O)	_	212		AIR		.127	.235	0.034
1 2 0	Greatte	• -	232		AIR	•	691.	. 269	0.038
130	MoS, +Pb0		316		AIR		.113	.200	0.061
) }	7								
1 3 1	MOS.+Moo.+Sh.O.	100	316	DRY	AIR	ŧ	.067	. 162	0.036
1 2 2	MOS-1003-0-2-4		316	DRY	AIR	:	. 134	.226	0.096
1 2 2	E 2 1 2 2 3 3		316		AIR		, 100	. 173	0.071
125	MOS-tono		316		AIR		.135	.202	0.043
* 0 1	10022 t 20 2 0 4) (127	1116	0.423
135	MoS2+T102	100	316	DKY	AIK		161.	117.	•
126	No.C. +CaO	100	316	DRY	AIR	ŧ	.046	.155	0.072
ם ה	2012 COURT	001	316		AIR		.089	. 154	0.109
2	ECZECTION OF ONE	100	316		AIR		180.	.169	0.077
06.1	Most shoot state of the state o	001	316		AIR		.095	. 188	0.067
77		001	274		AIR	•	.133	.195	0.061
•	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		•			٠			
1 7 1	Graphite	100	22	DRY	AIR	•	.113	.133	0.082
142		100	274	DRY	AIR	t	.059	.196	0.125
777		001	22	DRY	AIR		.165	. 196	0.113
			1 6	Adu	A 1 A		.167	.209	0.224
*		201	77	2 2	118		001	157	0.070
145			910				7	. 7 6	\$17.0
9 🤄 1	A1203+HoS2	100	316	DRY	AIR		911.	767.	7.0